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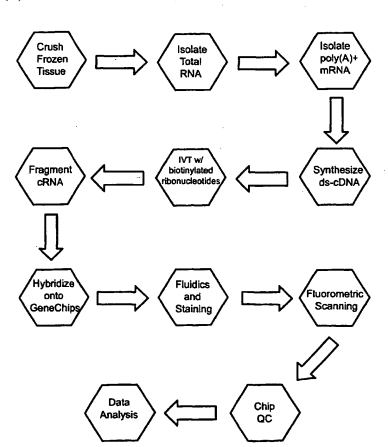
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(54) Title: GENE EXPRESSION PROFILES IN LIVER CANCER



(57) Abstract: The present invention identifies the global changes in gene expression associated with liver cancer by examining gene expression in tissue from normal liver, metastatic malignant liver and hepatocellular carcinoma. The present invention also identifies expression profiles which serve as useful diagnostic markers as well as markers that can be used to monitor disease states, disease progession, drug toxicity, druge efficacy and drug metabolism.



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GENE EXPRESSION PROFILES IN LIVER CANCER

INVENTORS

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RELATED APPLICATIONS

This application is related to U.S. Provisional Application 60/211,379, filed on June 14, 2000, and is related to U.S. Provisional Application 60/237,054, filed October 2, 2000, and is related to U.S. Application 09/880,107, filed June 14, 2001, each of which is herein incorporated by reference in its entirety.

BACKGROUND OF THE INVENTION

Primary hepatocellular carcinoma (HCC) is a widespread cancer throughout the world, especially prevalent where the incidence of chronic hepatitis B (HBV) and hepatitis C (HCV) viral infections are endemic (Groen, (1999) Semin. Oncol. Nurs. 15, 48-57; Idilman et al., (1998) J. Viral. Hepat. 5, 110-117; Di Bisceglie et al., (1998) Hepatol. 28, 1161-1165; Johnson, (1997) Hepatogastroenerology 44, 307-312; Sheu, (1997) J. Gastroeneterol. Hepatol. 12, S309-313). Hepatocellular carcinomas are very malignant tumors that generally offer a poor prognosis, dependent on the size of the tumor, the effect on normal liver functions, and the involvement of metastases. They are best treated by surgical resection, when the tumors are diagnosed at a stage where this is a viable possibility, but the recurrence rate for these cancers remains high (Johnson, (1997) Hepatogastroenterology 44, 307-312; Schafer & Sorrell, (1999) Lancet 353, 1253-1257; Groen, (1999) Semin. Oncol. Nurs. 15, 48-57; Sitzman, (1995) World. J. Surg. 19, 790-794; 20 DiCarlo, (1995) Hepato-Gastroenterol. 42, 222-259; Tanaka et al., (1996) Hepato-Gastroenterol. 43, 1172-1181; El-Assal et al., (1997) Surgery 122, 571-577).

Numerous risk factors for the development of HCC have been identified: cirrhosis, HBV or HCV infection, being male, alcohol-related liver disease, exposure to aflatoxins, vinyl chloride and radioactive thorium dioxide, cigarette smoking, ingestion of inorganic arsenic, the use of oral contraceptives and anabolic steroids, iron accumulation, and various inherited metabolic disorders (hemochromatosis, glycogen storage disease, porphyria,

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tyrosinemia, α-1-antitrypsin deficiency) (Di Bisceglie *et al.*, (1998) Hepatol. 28, 1161-1165; Chen *et al.*, (1997) J. Gastroenterol. Hepatol. 12, S294-308; Schafer & Sorrell (1999) Lancet 353, 1253-1257; Groen, (1999) Semin. Oncol. Nurs. 15, 48-57; Idilman *et al.*, (1998) J. Viral. Hepat. 5, 110-117; Johnson, (1997) Hepato-Gastroenterol. 44, 307-312).

In addition to liver tumors attributed to hepatocellular carcinoma, there are liver tumors that arise as metastases from primary tumors in other parts of the body. These tumors most often metastasize from the gastrointestinal organs, primarily the colon and rectum, but it is possible for metastatic liver cancers to occur from primary cancers throughout the body (Sitzman, (1990) Hepatic Neoplasia, in Bayless (editor) Current Therapy in Gastroenterology and Liver Disease, Marcel Dekker; Groen, (1999) Semin. Oncol. Nurs. 15, 48-57). These cancers can be treated using the routine therapies such as chemotherapy, radiotherapy, surgical resection, liver transplantation, chemoembolization, cryosurgery, or a combination of therapies (Sitzman 1990, Groen 1999).

The characterization of genes that are differentially expressed in tumorigenesis is an important step in identifying those that are intimately involved in the details of a cell's 15 transformation from normal to cancerous. Little is known about the molecular changes that take place in the liver during the course of tumor progression. While changes in the expression level of individual genes has been reported, for example, galectin-3 (Hsu et al., (1999) Int. J. Cancer 81: 519-526; Iurisci et al., (2000) Clin. Cancer Res. 6: 1389-1393; Nakamura et al., (1999) Int. J. Oncol. 15: 143-148) and pancreatic secretory trypsin 20 inhibitor (PTSI, Ohmachi et al., (1994) Int. J. Cancer 55: 728-734) the investigation of the global changes in gene expression which occur in liver have not been documented. The identification of genes that are expressed in tumor tissue at differing levels, but not expressed at any level in normal liver tissue will be very valuable for monitoring tumor progression. The identification of genes and ESTs that are expressed in both types of 25 tumors, i. e., primary hepatocellular carcinomas as well as metastatic tumors of a different origin, and not in normal liver cells would be extremely valuable for the diagnosis of liver cancer. Thus there exists a need in the art for the identification of new genes and ESTs to serve as molecular markers to monitor the onset and development of liver cancer. These and other needs are met by the present invention. 30

SUMMARY OF THE INVENTION

The present invention identifies the global changes in gene expression associated with liver cancer by examining gene expression in tissue from normal liver, metastatic malignant liver and hepatocellular carcinoma. The present invention also identifies expression profiles which serve as useful diagnostic markers as well as markers that can be used to monitor disease states, disease progression, drug toxicity, drug efficacy and drug metabolism.

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The invention includes methods of diagnosing the presence or absence of liver cancer in a patient comprising the step of detecting the level of expression in a tissue sample of two or more genes from Tables 3-9; wherein differential expression of the genes in Tables 3-9 is indicative of liver cancer. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3-5. In some preferred embodiments, the method may include detecting the expression level of one or more genes selected from a group consisting Tetraspan NET-6 protein; collagen, type V, alpha; and glypican 3.

The invention also includes methods of detecting the progression of liver cancer and/or differentiating nonmetastatic from metastatic disease. For instance, methods of the invention include detecting the progression of liver cancer in a patient comprising the step of detecting the level of expression in a tissue sample of two or more genes from 3-9; wherein differential expression of the genes in Tables 3-9 is indicative of liver cancer progression. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3-5.

In some aspects, the present invention provides a method of monitoring the treatment of a patient with liver cancer, comprising administering a pharmaceutical composition to the patient and preparing a gene expression profile from a cell or tissue sample from the patient and comparing the patient gene expression profile to a gene expression from a cell population comprising normal liver cells or to a gene expression profile from a cell population comprising liver cancer cells or to both. In some preferred embodiments, the gene profile will include the expression level of one or more genes in Tables 3-9. In other preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3-5.

In another aspect, the present invention provides a method of treating a patient with liver cancer, comprising administering to the patient a pharmaceutical composition, wherein

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the composition alters the expression of at least one gene in Tables 3-9, preparing a gene expression profile from a cell or tissue sample from the patient comprising tumor cells and comparing the patient expression profile to a gene expression profile from an untreated cell population comprising liver cancer cells. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3-5.

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In one aspect, the present invention provides a method of diagnosing hepatocellular carcinoma in a patient, comprising detecting the level of expression in a tissue sample of two or more genes from Tables 3-9, wherein differential expression of the genes in Tables 3-9 is indicative of hepatocellular carcinoma. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3 or 5.

In another aspect, the present invention provides a method of detecting the progression of hepatocellular carcinoma in a patient, comprising detecting the level of expression in a tissue sample of two or more genes from Tables 3-9; wherein differential expression of the genes in Tables 3-9 is indicative of hepatocellular carcinoma progression. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3 or 5.

The present invention also provides materials and methods for monitoring the treatment of a patient with a hepatocellular caricnoma. The present invention provides a method of monitoring the treatment of a patient with hepatocellular carcinoma, comprising administering a pharmaceutical composition to the patient, preparing a gene expression profile from a cell or tissue sample from the patient and comparing the patient gene expression profile to a gene expression from a cell population comprising normal liver cells or to a gene expression profile from a cell population comprising hepatocellular carcinoma cells or to both. In some preferred embodiments, the method may include detecting the level of expression of one or more genes from the genes listed in Tables 3-9. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3 or 5.

In a related aspect, the present invention provides a method of treating a patient with hepatocellular carcinoma, comprising administering to the patient a pharmaceutical composition, wherein the composition alters the expression of at least one gene in Tables 3-9, preparing a gene expression profile from a cell or tissue sample from the patient comprising hepatocellular carcinoma cells and comparing the patient expression profile to a

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gene expression profile from an untreated cell population comprising hepatocellular carcinoma cells. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3 or 5.

The present invention provides a method of diagnosing a metastatic liver tumor in a patient, comprising detecting the level of expression in a tissue sample of two or more genes from Tables 3-9, wherein differential expression of the genes in Tables 3-9 is indicative of a metastatic liver tumor. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 4 or 5.

The present invention provides a method of detecting the progression of a metastatic liver tumor in a patient, comprising detecting the level of expression in a tissue sample of two or more genes from Tables 3-9, wherein differential expression of the genes in Tables 3-9 is indicative of a metastatic liver tumor progression. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 4 or 5.

In a related aspect, the present invention provides a method of monitoring the treatment of a patient with a metastatic liver tumor, comprising administering a pharmaceutical composition to the patient, preparing a gene expression profile from a cell or tissue sample from the patient and comparing the patient gene expression profile to a gene expression from a cell population comprising normal liver cells or to a gene expression profile from a cell population comprising metastatic liver tumor cells or to both. In some preferred embodiments, the method of the present invention may include detecting the expression level of one or more genes selected from the genes listed in Tables 3-9. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 4 or 5.

In some preferred embodiments, the present invention provides a method of treating a patient with a metastatic liver tumor, comprising administering to the patient a pharmaceutical composition, wherein the composition alters the expression of at least one gene in Tables 3-9, preparing a gene expression profile from a cell or tissue sample from the patient comprising metastatic liver tumor cells and comparing the patient expression profile to a gene expression profile from an untreated cell population comprising metastatic liver tumor cells. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 4 or 5.

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The invention also includes methods of differentiating metastatic liver cancer from hepatocellular carcinoma in a patient comprising the step of detecting the level of expression in a tissue sample of two or more genes from Tables 3-9; wherein differential expression of the genes in Tables 3-9 is indicative of metastatic liver cancer rather than hepatocellular carcinoma.

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The invention further includes methods of screening for an agent capable of modulating the onset or progression of liver cancer, comprising the steps of exposing a cell to the agent; and detecting the expression level of two or more genes from Tables 3-9. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3-5.

Any of the methods of the invention described above may include the detection of at least 2 genes from the tables. Preferred methods may detect all or nearly all of the genes in the tables. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3-5.

The invention further includes compositions comprising at least two oligonucleotides, wherein each of the oligonucleotides comprises a sequence that specifically hybridizes to a gene in Tables 3-9 as well as solid supports comprising at least two probes, wherein each of the probes comprises a sequence that specifically hybridizes to a gene in Tables 3-9. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3-5.

The invention further includes computer systems comprising a database containing information identifying the expression level in liver tissue of a set of genes comprising at least two genes in Tables 3-9; and a user interface to view the information. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3-5. The database may further include sequence information for the genes, information identifying the expression level for the set of genes in normal liver tissue and malignant tissue (metastatic and nonmetastatic) and may contain links to external databases such as GenBank.

Lastly, the invention includes methods of using the databases, such as methods of using the disclosed computer systems to present information identifying the expression level in a tissue or cell of at least one gene in Tables 3-9, comprising the step of comparing the expression level of at least one gene in Tables 3-9 in the tissue or cell to the level of

expression of the gene in the database. In some preferred embodiments, one or more genes may be selected from a group consisting of the genes listed in Tables 3-5.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a flow chart showing a schematic representation of the experimental protocol.

Figures 2A-2C are graphs of the number of genes present in all samples as a function of the number of samples for the second sample set.

10 DETAILED DESCRIPTION

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Many biological functions are accomplished by altering the expression of various genes through transcriptional (e.g., through control of initiation, provision of RNA precursors, RNA processing, etc.) and/or translational control. For example, fundamental biological processes such as cell cycle, cell differentiation and cell death, are often characterized by the variations in the expression levels of groups of genes.

Changes in gene expression also are associated with pathogenesis. For example, the lack of sufficient expression of functional tumor suppressor genes and/or the over expression of oncogene/protooncogenes could lead to tumorgenesis or hyperplastic growth of cells (Marshall, (1991) Cell, 64, 313-326; Weinberg, (1991) Science, 254, 1138-1146). Thus, changes in the expression levels of particular genes (e.g., oncogenes or tumor suppressors) serve as signposts for the presence and progression of various diseases.

Monitoring changes in gene expression may also provide certain advantages during drug screening development. Often drugs are screened and prescreened for the ability to interact with a major target without regard to other effects the drugs have on cells. Often such other effects cause toxicity in the whole animal, which prevent the development and use of the potential drug.

The present inventors have examined tissue samples from normal liver, metastatic malignant liver and hepatocellular carcinoma to identify the global changes in gene expression associated with liver cancer. The protocol used is schematically represented in Figure 1. These global changes in gene expression, also referred to as expression profiles, provide useful markers for diagnostic uses as well as markers that can be used to monitor disease states, disease progression, drug toxicity, drug efficacy and drug metabolism.

Definitions

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In the description that follows, numerous terms and phrases known to those skilled in the art are used. In the interests of clarity and consistency of interpretation, the definitions of certain terms and phrases are provided.

The present invention provides compositions and methods to detect the level of expression of genes that may be differentially expressed dependent upon the state of the cell, *i.e.*, normal versus cancerous. As used herein, the phrase "detecting the level of expression" includes methods that quantitate expression levels as well as methods that determine whether a gene of interest is expressed at all. Thus, an assay which provides a yes or no result without necessarily providing quantification of an amount of expression is an assay that requires "detecting the level of expression" as that phrase is used herein.

As used herein, oligonucleotide sequences that are complementary to one or more of the genes described herein, refers to oligonucleotides that are capable of hybridizing under stringent conditions to at least part of the nucleotide sequence of said genes. Such hybridizable oligonucleotides will typically exhibit at least about 75% sequence identity at the nucleotide level to said genes, preferably about 80% or 85% sequence identity or more preferably about 90% or 95% or more sequence identity to said genes.

"Bind(s) substantially" refers to complementary hybridization between a probe nucleic acid and a target nucleic acid and embraces minor mismatches that can be accommodated by reducing the stringency of the hybridization media to achieve the desired detection of the target polynucleotide sequence.

The terms "background" or "background signal intensity" refer to hybridization signals resulting from non-specific binding, or other interactions, between the labeled target nucleic acids and components of the oligonucleotide array (e.g., the oligonucleotide probes, control probes, the array substrate, etc.). Background signals may also be produced by intrinsic fluorescence of the array components themselves. A single background signal can be calculated for the entire array, or a different background signal may be calculated for each target nucleic acid. In a preferred embodiment, background is calculated as the average hybridization signal intensity for the lowest 5% to 10% of the probes in the array, or, where a different background signal is calculated for each target gene, for the lowest 5% to 10% of the probes for each gene. Of course, one of skill in the art will appreciate that

where the probes to a particular gene hybridize well and thus appear to be specifically binding to a target sequence, they should not be used in a background signal calculation. Alternatively, background may be calculated as the average hybridization signal intensity produced by hybridization to probes that are not complementary to any sequence found in the sample (e.g., probes directed to nucleic acids of the opposite sense or to genes not found in the sample such as bacterial genes where the sample is mammalian nucleic acids). Background can also be calculated as the average signal intensity produced by regions of the array that lack any probes at all.

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The phrase "hybridizing specifically to" refers to the binding, duplexing or hybridizing of a molecule substantially to or only to a particular nucleotide sequence or sequences under stringent conditions when that sequence is present in a complex mixture (e.g., total cellular) DNA or RNA.

The term "mismatch control" or "mismatch probe" refer to a probe whose sequence is deliberately selected not to be perfectly complementary to a particular target sequence. For each mismatch (MM) control in a high-density array there typically exists a corresponding perfect match (PM) probe that is perfectly complementary to the same particular target sequence. The mismatch may comprise one or more bases.

While the mismatch(s) may be located anywhere in the mismatch probe, terminal mismatches are less desirable as a terminal mismatch is less likely to prevent hybridization of the target sequence. In a particularly preferred embodiment, the mismatch is located at or near the center of the probe such that the mismatch is most likely to destabilize the duplex with the target sequence under the test hybridization conditions.

The term "perfect match probe" refers to a probe that has a sequence that is perfectly complementary to a particular target sequence. The test probe is typically perfectly complementary to a portion (subsequence) of the target sequence. The perfect match (PM) probe can be a "test probe", a "normalization control" probe, an expression level control probe and the like. A perfect match control or perfect match probe is, however, distinguished from a "mismatch control" or "mismatch probe."

As used herein a "probe" is defined as a nucleic acid, capable of binding to a target nucleic acid of complementary sequence through one or more types of chemical bonds, usually through complementary base pairing, usually through hydrogen bond formation. As used herein, a probe may include natural (i.e., A, G, U, C or T) or modified bases (7-

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deazaguanosine, inosine, etc.). In addition, the bases in probes may be joined by a linkage other than a phosphodiester bond, so long as it does not interfere with hybridization. Thus, probes may be peptide nucleic acids in which the constituent bases are joined by peptide bonds rather than phosphodiester linkages.

The term "stringent conditions" refers to conditions under which a probe will hybridize to its target subsequence, but with only insubstantial hybridization to other sequences or to other sequences such that the difference may be identified. Stringent conditions are sequence-dependent and will be different in different circumstances. Longer sequences hybridize specifically at higher temperatures. Generally, stringent conditions are selected to be about 5°C lower than the thermal melting point (T_m) for the specific sequence at a defined ionic strength and pH.

Typically, stringent conditions will be those in which the salt concentration is at least about 0.01 to 1.0 M sodium ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes (e.g., 10 to 50 nucleotide). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide.

The "percentage of sequence identity" or "sequence identity" is determined by comparing two optimally aligned sequences or subsequences over a comparison window or span, wherein the portion of the polynucleotide sequence in the comparison window may optionally comprise additions or deletions (*i.e.*, gaps) as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at which the identical monomer unit (*e.g.*, nucleic acid base or amino acid residue) occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the window of comparison and multiplying the result by 100 to yield the percentage of sequence identity. Percentage sequence identity when calculated using the programs GAP or BESTFIT (see below) is calculated using default gap weights.

Homology or identity may be determined by BLAST (Basic Local Alignment

Search Tool) analysis using the algorithm employed by the programs blastp, blastn, blastx,
tblastn and tblastx (Karlin et al., (1990) Proc. Natl. Acad. Sci. USA 87, 2264-2268 and
Altschul, (1993) J. Mol. Evol. 36, 290-300, fully incorporated by reference) which are

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tailored for sequence similarity searching. The approach used by the BLAST program is to first consider similar segments between a query sequence and a database sequence, then to evaluate the statistical significance of all matches that are identified and finally to summarize only those matches which satisfy a preselected threshold of significance. For a discussion of basic issues in similarity searching of sequence databases, see Altschul et al., 5 (1994) Nature Genet. 6, 119-129) which is fully incorporated by reference. The search parameters for histogram, descriptions, alignments, expect (i.e., the statistical significance threshold for reporting matches against database sequences), cutoff, matrix and filter are at the default settings. The default scoring matrix used by blastp, blastx, tblastn, and tblastx is the BLOSUM62 matrix (Henikoff et al., (1992) Proc. Natl. Acad. Sci. USA 89, 10915-10 10919, fully incorporated by reference). Four blastn parameters were adjusted as follows: Q=10 (gap creation penalty); R=10 (gap extension penalty); wink=1 (generates word hits at every winkth position along the query); and gapw=16 (sets the window width within which gapped alignments are generated). The equivalent Blastp parameter settings were Q=9; R=2; wink=1; and gapw=32. A Bestfit comparison between sequences, available in the 15 GCG package version 10.0, uses DNA parameters GAP=50 (gap creation penalty) and LEN=3 (gap extension penalty) and the equivalent settings in protein comparisons are GAP=8 and LEN=2.

Uses for the Liver Cancer Markers as Diagnostics 20

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As described herein, the genes and gene expression information provided in Tables 3-9 may be used as diagnostic markers for the prediction or identification of the malignant state of the liver tissue. For instance, a liver tissue sample or other sample from a patient may be assayed by any of the methods described herein or by any other method known to those skilled in the art, and the expression levels from a gene or genes from the Tables, in particular the genes in Tables 3-5, may be compared to the expression levels found in normal liver tissue, tissue from metastatic liver cancer or hepatocellular carcinoma tissue. Expression profiles generated from the tissue or other sample that substantially resemble an expression profile from normal or diseased liver tissue may be used, for instance, to aid in disease diagnosis. Comparison of the expression data, as well as available sequence or other information may be done by researcher or diagnostician or may be done with the aid of a computer and databases as described above.

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Use of the Liver Cancer Markers for Monitoring Disease Progression

As described above, the genes and gene expression information provided in Tables 3-9 may also be used as markers for the monitoring of disease progression, for instance, the development of liver cancer. For instance, a liver tissue sample or other sample from a patient may be assayed by any of the methods described above, and the expression levels in the sample from a gene or genes from or 3-9 may be compared to the expression levels found in normal liver tissue, tissue from metastatic liver cancer or hepatocellular carcinoma tissue. Comparison of the expression data, as well as available sequence or other information may be done by researcher or diagnostician or may be done with the aid of a computer and databases as described above.

Use of the Liver Cancer Markers for Drug Screening

According to the present invention, the genes identified in Tables 3-9 may be used as markers to evaluate the effects of a candidate drug or agent on a cell, particularly a cell undergoing malignant transformation, for instance, a liver cancer cell or tissue sample. A candidate drug or agent can be screened for the ability to stimulate the transcription or expression of a given marker or markers (drug targets) or to down-regulate or counteract the transcription or expression of a marker or markers. According to the present invention, one can also compare the specificity of drugs' effects by looking at the number of markers affected by different drugs and comparing them. More specific drugs will affect fewer transcriptional targets. Similar sets of markers identified for two drugs indicates a similarity of effects.

Agents that are assayed in the above methods can be randomly selected or rationally selected or designed. As used herein, an agent is said to be randomly selected when the agent is chosen randomly without considering the specific sequences involved in the association of the a protein of the invention alone or with its associated substrates, binding partners, etc. An example of randomly selected agents is the use a chemical library or a peptide combinatorial library, or a growth broth of an organism.

As used herein, an agent is said to be rationally selected or designed when the agent is chosen on a nonrandom basis which takes into account the sequence of the target site and/or its conformation in connection with the agents action. Agents can be rationally

selected or rationally designed by utilizing the peptide sequences that make up these sites. For example, a rationally selected peptide agent can be a peptide whose amino acid sequence is identical to or a derivative of any functional consensus site.

The agents of the present invention can be, as examples, peptides, small molecules, vitamin derivatives, as well as carbohydrates. Dominant negative proteins, DNA encoding these proteins, antibodies to these proteins, peptide fragments of these proteins or mimics of these proteins may be introduced into cells to affect function. "Mimic" as used herein refers to the modification of a region or several regions of a peptide molecule to provide a structure chemically different from the parent peptide but topographically and functionally similar to the parent peptide (see Grant, (1995) in Molecular Biology and Biotechnology Meyers (editor) VCH Publishers). A skilled artisan can readily recognize that there is no limit as to the structural nature of the agents of the present invention.

Assay Formats

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The genes identified as being differentially expressed in liver cancer may be used in a variety of nucleic acid detection assays to detect or quantititate the expression level of a gene or multiple genes in a given sample. Any hybridization assay format may be used, including solution-based and solid support-based assay formats, for example, traditional Northern blotting. Other suitable assay formats that may be used for detecting gene expression levels include, but are not limited to, nuclease protection, RT-PCR and differential display methods. These methods are useful for some embodiments of the invention; however, methods and assays of the invention are most efficiently designed with array or chip hybridization-based methods for detecting the expression of a large number of genes. Assays and methods of the invention may utilize available formats to simultaneously screen at least about 100, preferably about 1000, more preferably about 10,000 and most preferably about 1,000,000 or more different nucleic acid hybridizations.

Assays to monitor the expression of a marker or markers as defined in Tables 3-9 may utilize any available means of monitoring for changes in the expression level of the nucleic acids of the invention. As used herein, an agent is said to modulate the expression of a nucleic acid of the invention if it is capable of up- or down-regulating expression of the nucleic acid in a cell.

In one assay format, gene chips containing probes to at least two genes from Tables

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3-9 may be used to directly monitor or detect changes in gene expression in the treated or exposed cell as described in more detail above. In another format, cell lines that contain reporter gene fusions between the open reading frame and/or the 3' or 5' regulatory regions of a gene in Tables 3-9 and any assayable fusion partner may be prepared. Numerous assayable fusion partners are known and readily available including the firefly luciferase gene and the gene encoding chloramphenical acetyltransferase (Alam *et al.*, (1990) Anal. Biochem. 188, 245-254). Cell lines containing the reporter gene fusions are then exposed to the agent to be tested under appropriate conditions and time. Differential expression of the reporter gene between samples exposed to the agent and control samples identifies agents which modulate the expression of the nucleic acid.

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Additional assay formats may be used to monitor the ability of the agent to modulate the expression of one or more genes identified in Tables 3-9. For instance, as described above, mRNA expression may be monitored directly by hybridization of probes to the nucleic acids of the invention. Cell lines are exposed to the agent to be tested under appropriate conditions and time and total RNA or mRNA is isolated by standard procedures such those disclosed in Sambrook *et al.*, (1989) Molecular Cloning - A Laboratory Manual, Cold Spring Harbor Laboratory Press).

In another assay format, cells or cell lines are first identified which express the gene products of the invention physiologically. Cell and/or cell lines so identified would be expected to comprise the necessary cellular machinery such that the fidelity of modulation of the transcriptional apparatus is maintained with regard to exogenous contact of agent with appropriate surface transduction mechanisms and/or the cytosolic cascades. Such cell lines may be, but are not required to be, derived from liver tissue. Further, such cells or cell lines may be transduced or transfected with an expression vehicle (e.g., a plasmid or viral vector) construct comprising an operable non-translated 5'-promoter containing end of the structural gene encoding the instant gene products fused to one or more antigenic fragments, which are peculiar to the instant gene products, wherein said fragments are under the transcriptional control of said promoter and are expressed as polypeptides whose molecular weight can be distinguished from the naturally occurring polypeptides or may further comprise an immunologically distinct tag. Such a process is well known in the art (see Sambrook et al., (1989) Molecular Cloning - A Laboratory Manual, Cold Spring Harbor Laboratory Press).

Cells or cell lines transduced or transfected as outlined above are then contacted with agents under appropriate conditions; for example, the agent comprises a pharmaceutically acceptable excipient and is contacted with cells comprised in an aqueous physiological buffer such as phosphate buffered saline (PBS) at physiological pH, Eagles balanced salt solution (BSS) at physiological pH, PBS or BSS comprising serum or conditioned media comprising PBS or BSS and serum incubated at 37°C. Said conditions may be modulated as deemed necessary by one of skill in the art. Subsequent to contacting the cells with the agent, said cells will be disrupted and the polypeptides of the lysate are fractionated such that a polypeptide fraction is pooled and contacted with an antibody to be further processed by immunological assay (e.g., ELISA, immunoprecipitation or Western blot). The pool of proteins isolated from the "agent-contacted" sample will be compared with a control sample where only the excipient is contacted with the cells and an increase or decrease in the immunologically generated signal from the "agent-contacted" sample compared to the control will be used to distinguish the effectiveness of the agent.

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Another embodiment of the present invention provides methods for identifying agents that modulate the levels, concentration or at least one activity of a protein(s) encoded by the genes in Tables 3-9. Such methods or assays may utilize any means of monitoring or detecting the desired activity.

In one format, the relative amounts of a protein of the invention between a cell population that has been exposed to the agent to be tested compared to an un-exposed control cell population may be assayed. In this format, probes such as specific antibodies are used to monitor the differential expression of the protein in the different cell populations. Cell lines or populations are exposed to the agent to be tested under appropriate conditions and time. Cellular lysates may be prepared from the exposed cell line or population and a control, unexposed cell line or population. The cellular lysates are then analyzed with the probe, such as a specific antibody.

The genes which are assayed according to the present invention are typically in the form of mRNA or reverse transcribed mRNA. The genes may be cloned or not and the genes may be amplified or not. The cloning itself does not appear to bias the representation of genes within a population. However, it may be preferable to use polyA+ RNA as a source, as it can be used with less processing steps.

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The sequences of the expression marker genes are in the public databases. Tables 3-9 provide the Affymetrix gene ID and GenBank accession number for each marker identified. The nucleotide sequence for each marker has been provided in electronic format with this application and these sequences are incorporated herein by reference as are equivalent and related sequences present in the public databases.

Probes based on the sequences of the genes described herein may be prepared by any commonly available method. Oligonucleotide probes for assaying the tissue or cell sample are preferably of sufficient length to specifically hybridize only to appropriate, complementary genes or transcripts. Typically the oligonucleotide probes will be at least 10, 12, 14, 16, 18, 20 or 25 nucleotides in length. In some cases longer probes of at least 30, 40, or 50 nucleotides will be desirable.

Probe design

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One of skill in the art will appreciate that an enormous number of array designs are suitable for the practice of this invention. The high density array will typically include a number of probes that specifically hybridize to the sequences of interest. See WO 99/32660 for methods of producing probes for a given gene or genes. In addition, in a preferred embodiment, the array will include one or more control probes.

High density array chips of the invention include "test probes." Test probes may be oligonucleotides that range from about 5 to about 500 or about 50 nucleotides, more preferably from about 10 to about 40 nucleotides and most preferably from about 15 to about 40 nucleotides in length. In other particularly preferred embodiments the probes are about 20 or 25 nucleotides in length. In another preferred embodiment, test probes are double or single strand DNA sequences. DNA sequences are isolated or cloned from natural sources or amplified from natural sources using natural nucleic acid as templates. These probes have sequences complementary to particular subsequences of the genes whose expression they are designed to detect. Thus, the test probes are capable of specifically hybridizing to the target nucleic acid they are to detect.

In addition to test probes that bind the target nucleic acid(s) of interest, the high density array can contain a number of control probes. The control probes fall into three categories referred to herein as (1) normalization controls; (2) expression level controls; and (3) mismatch controls.

Normalization controls are oligonucleotide or other nucleic acid probes that are complementary to labeled reference oligonucleotides or other nucleic acid sequences that are added to the nucleic acid sample. The signals obtained from the normalization controls after hybridization provide a control for variations in hybridization conditions, label intensity, "reading" efficiency and other factors that may cause the signal of a perfect hybridization to vary between arrays. In a preferred embodiment, signals (e.g., fluorescence intensity) read from all other probes in the array are divided by the signal (e.g., fluorescence intensity) from the control probes thereby normalizing the measurements.

Virtually any probe may serve as a normalization control. However, it is recognized that hybridization efficiency varies with base composition and probe length. Preferred normalization probes are selected to reflect the average length of the other probes present in the array, however, they can be selected to cover a range of lengths. The normalization control(s) can also be selected to reflect the (average) base composition of the other probes in the array, however in a preferred embodiment, only one or a few probes are used and they are selected such that they hybridize well (i.e., no secondary structure) and do not match any target-specific probes.

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Expression level controls are probes that hybridize specifically with constitutively expressed genes in the biological sample. Virtually any constitutively expressed gene provides a suitable target for expression level controls. Typical expression level control probes have sequences complementary to subsequences of constitutively expressed "housekeeping genes" including, but not limited to the β-actin gene, the transferrin receptor gene, the GAPDH gene, and the like.

Mismatch controls may also be provided for the probes to the target genes, for expression level controls or for normalization controls. Mismatch controls are oligonucleotide probes or other nucleic acid probes identical to their corresponding test or control probes except for the presence of one or more mismatched bases. A mismatched base is a base selected so that it is not complementary to the corresponding base in the target sequence to which the probe would otherwise specifically hybridize. One or more mismatches are selected such that under appropriate hybridization conditions (e.g., stringent conditions) the test or control probe would be expected to hybridize with its target sequence, but the mismatch probe would not hybridize (or would hybridize to a significantly lesser extent). Preferred mismatch probes contain a central mismatch. Thus, for example, where a

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probe is a twenty-mer, a corresponding mismatch probe will have the identical sequence except for a single base mismatch (e.g., substituting a G, a C or a T for an A) at any of positions 6 through 14 (the central mismatch).

Mismatch probes thus provide a control for non-specific binding or cross hybridization to a nucleic acid in the sample other than the target to which the probe is directed. Mismatch probes also indicate whether a hybridization is specific or not. For example, if the target is present the perfect match probes should be consistently brighter than the mismatch probes. In addition, if all central mismatches are present, the mismatch probes can be used to detect a mutation. The difference in intensity between the perfect match and the mismatch probe (I(PM) - I(MM)) provides a good measure of the concentration of the hybridized material.

Nucleic Acid Samples

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As is apparent to one of ordinary skill in the art, nucleic acid samples used in the methods and assays of the invention may be prepared by any available method or process. Methods of isolating total mRNA are also well known to those of skill in the art. For example, methods of isolation and purification of nucleic acids are described in detail in Chapter 3 of Laboratory Techniques in Biochemistry and Molecular Biology: Hybridization With Nucleic Acid Probes, Part I Theory and Nucleic Acid Preparation, Tijssen, (1993) (editor) Elsevier Press. Such samples include RNA samples, but also include cDNA synthesized from a mRNA sample isolated from a cell or tissue of interest. Such samples also include DNA amplified from the cDNA, and an RNA transcribed from the amplified DNA. One of skill in the art would appreciate that it is desirable to inhibit or destroy RNase present in homogenates before homogenates can be used.

Biological samples may be of any biological tissue or fluid or cells from any organism as well as cells raised *in vitro*, such as cell lines and tissue culture cells. Frequently the sample will be a "clinical sample" which is a sample derived from a patient. Typical clinical samples include, but are not limited to, sputum, blood, blood-cells (e.g., white cells), tissue or fine needle biopsy samples, urine, peritoneal fluid, and pleural fluid, or cells therefrom.

Biological samples may also include sections of tissues, such as frozen sections or formalin fixed sections taken for histological purposes.

Solid Supports

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Solid supports containing oligonucleotide probes for differentially expressed genes of the invention can be filters, polyvinyl chloride dishes, silicon or glass based chips, etc. Such wafers and hybridization methods are widely available, for example, those disclosed by Beattie (WO 95/11755). Any solid surface to which oligonucleotides can be bound, either directly or indirectly, either covalently or non-covalently, can be used. A preferred solid support is a high density array or DNA chip. These contain a particular oligonucleotide probe in a predetermined location on the array. Each predetermined location may contain more than one molecule of the probe, but each molecule within the predetermined location has an identical sequence. Such predetermined locations are termed features. There may be, for example, about 2, 10, 100, 1000 to 10,000; 100,000 or 400,000 of such features on a single solid support. The solid support, or the area within which the probes are attached may be on the order of a square centimeter.

Oligonucleotide probe arrays for expression monitoring can be made and used according to any techniques known in the art (see for example, Lockhart *et al.*, (1996) Nat. Biotechnol. 14, 1675-1680; McGall *et al.*, (1996) Proc. Nat. Acad. Sci. USA 93, 13555-13460). Such probe arrays may contain at least two or more oligonucleotides that are complementary to or hybridize to two or more of the genes described herein. Such arrays may also contain oligonucleotides that are complementary or hybridize to at least about 2, 3, 4, 5, 6, 7, 8, 9, 10, 20, 30, 50, 70, 100 or or more the genes described herein.

Methods of forming high density arrays of oligonucleotides with a minimal number of synthetic steps are known. The oligonucleotide analogue array can be synthesized on a solid substrate by a variety of methods, including, but not limited to, light-directed chemical coupling, and mechanically directed coupling (see Pirrung et al., (1992) U.S. Patent No. 5,143, 854; Fodor et al., (1998) U.S. Patent No. 5,800,992; Chee et al., (1998) 5,837,832.

In brief, the light-directed combinatorial synthesis of oligonucleotide arrays on a glass surface proceeds using automated phosphoramidite chemistry and chip masking techniques. In one specific implementation, a glass surface is derivatized with a silane reagent containing a functional group, e.g., a hydroxyl or amine group blocked by a photolabile protecting group. Photolysis through a photolithogaphic mask is used selectively to expose functional groups which are then ready to react with incoming 5'

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photoprotected nucleoside phosphoramidites. The phosphoramidites react only with those sites which are illuminated (and thus exposed by removal of the photolabile blocking group). Thus, the phosphoramidites only add to those areas selectively exposed from the preceding step. These steps are repeated until the desired array of sequences have been synthesized on the solid surface. Combinatorial synthesis of different oligonucleotide analogues at different locations on the array is determined by the pattern of illumination during synthesis and the order of addition of coupling reagents.

In addition to the foregoing, additional methods which can be used to generate an array of oligonucleotides on a single substrate are described in Fodor *et al.*, (1993). WO 93/09668. High density nucleic acid arrays can also be fabricated by depositing premade or natural nucleic acids in predetermined positions. Synthesized or natural nucleic acids are deposited on specific locations of a substrate by light directed targeting and oligonucleotide directed targeting. Another embodiment uses a dispenser that moves from region to region to deposit nucleic acids in specific spots.

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Hybridization

Nucleic acid hybridization simply involves contacting a probe and target nucleic acid under conditions where the probe and its complementary target can form stable hybrid duplexes through complementary base pairing (see Lockhart *et al.*, (1999) WO 99/32660). The nucleic acids that do not form hybrid duplexes are then washed away leaving the hybridized nucleic acids to be detected, typically through detection of an attached detectable label. It is generally recognized that nucleic acids are denatured by increasing the temperature or decreasing the salt concentration of the buffer containing the nucleic acids.

Under low stringency conditions (e.g., low temperature and/or high salt) hybrid duplexes (e.g., DNA-DNA, RNA-RNA or RNA-DNA) will form even where the annealed sequences are not perfectly complementary.

Thus specificity of hybridization is reduced at lower stringency. Conversely, at higher stringency (e.g., higher temperature or lower salt) successful hybridization requires fewer mismatches. One of skill in the art will appreciate that hybridization conditions may be selected to provide any degree of stringency. In a preferred embodiment, hybridization is performed at low stringency, in this case in 6× SSPE-T at 37°C (0.005% Triton x-100) to ensure hybridization and then subsequent washes are performed at higher stringency (e.g.,

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1× SSPE-T at 37°C) to eliminate mismatched hybrid duplexes. Successive washes may be performed at increasingly higher stringency (e.g., down to as low as 0.25× SSPET at 37°C to 50°C) until a desired level of hybridization specificity is obtained. Stringency can also be increased by addition of agents such as formamide. Hybridization specificity may be evaluated by comparison of hybridization to the test probes with hybridization to the various controls that can be present (e.g., expression level control, normalization control, mismatch controls, etc.).

In general, there is a tradeoff between hybridization specificity (stringency) and signal intensity. Thus, in a preferred embodiment, the wash is performed at the highest stringency that produces consistent results and that provides a signal intensity greater than approximately 10% of the background intensity. Thus, in a preferred embodiment, the hybridized array may be washed at successively higher stringency solutions and read between each wash. Analysis of the data sets thus produced will reveal a wash stringency above which the hybridization pattern is not appreciably altered and which provides adequate signal for the particular oligonucleotide probes of interest.

Signal Detection

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The hybridized nucleic acids are typically detected by detecting one or more labels attached to the sample nucleic acids. The labels may be incorporated by any of a number of means well known to those of skill in the art (see Lockhart *et al.*, (1999) WO 99/32660).

Databases

The present invention includes relational databases containing sequence information, for instance for the genes of Tables 3-9, as well as gene expression information in various liver tissue samples. Databases may also contain information associated with a given sequence or tissue sample such as descriptive information about the gene associated with the sequence information, or descriptive information concerning the clinical status of the tissue sample, or the patient from which the sample was derived. The database may be designed to include different parts, for instance a sequences database and a gene expression database. Methods for the configuration and construction of such databases are widely available, for instance, see Akerblom *et al.*, (1999) U.S. Patent 5,953,727, which is herein incorporated by reference in its entirety.

The databases of the invention may be linked to an outside or external database. In a preferred embodiment, as described in Tables 3-9, the external database is GenBank and the associated databases maintained by the National Center for Biotechnology Information (NCBI).

Any appropriate computer platform may be used to perform the necessary comparisons between sequence information, gene expression information and any other information in the database or provided as an input. For example, a large number of computer workstations are available from a variety of manufacturers, such has those available from Silicon Graphics. Client-server environments, database servers and networks are also widely available and appropriate platforms for the databases of the invention.

The databases of the invention may be used to produce, among other things, electronic Northerns to allow the user to determine the cell type or tissue in which a given gene is expressed and to allow determination of the abundance or expression level of a given gene in a particular tissue or cell.

The databases of the invention may also be used to present information identifying the expression level in a tissue or cell of a set of genes comprising at least one gene in Tables 3-9 comprising the step of comparing the expression level of at least one gene in Tables 3-9 in the tissue to the level of expression of the gene in the database. Such methods may be used to predict the physiological state of a given tissue by comparing the level of expression of a gene or genes in Tables 3-9 from a sample to the expression levels found in tissue from normal liver, malignant liver or hepatocellular carcinoma. Such methods may also be used in the drug or agent screening assays as described below.

Without further description, it is believed that one of ordinary skill in the art can, using the preceding description and the following illustrative examples, make and utilize the compounds of the present invention and practice the claimed methods. The following working examples therefore, specifically point out the preferred embodiments of the present invention, and are not to be construed as limiting in any way the remainder of the disclosure.

EXAMPLES

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Example 1: Tissue Sample Acquisition and Preparation 30

Figure 1 outlines the experimental protocol used. Liver tissue samples were excised and snap frozen in liquid nitrogen. The clinical data for each of the samples included in this WO 02/29103

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study are outlined in Table 1. The sample set was composed of eight samples of normal liver tissue (N1-N8), five samples of metastatic adenocarcinoma arising from rectum (designated M1 and M3) and colon (M2, M4 and M5) tissues and six samples of primary hepatocellular carcinomas. Samples were named according to type of tissue:

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HCC=hepatocellular carcinoma, M=metastatic, N=normal. Table 1 includes the TNM classification (the American Joint Committee on Cancer's system of classifying cancers) of the tissues used as samples where T refers to the extent of the primary tumor, N refers to the absence or presence and extent of regional lymph node metastasis, and M refers to the absence or presence of distant metastasis. Numbers following T, N, and M refer to the size of the primary tumor and the amount of vascular invasion, where 0=no evidence of tumor, lymph node involvement or metastasis, 4=multiple tumors involved, and x=cannot be assessed. Histopathologic grade (Table 1) is a qualitative assessment of differentiation of a tumor, where G1=most differentiated and G4=undifferentiated. Clinical stage (Table 1) characterizes the anatomic extent of disease in the patient from whom the sample was taken, where I and II are early stages, III and IV are late stages.

With minor modifications, the sample preparation protocol followed the Affymetrix GeneChip Expression Analysis Manual. Frozen tissue was first ground to powder using the Spex Certiprep 6800 Freezer Mill. Total RNA was then extracted using Trizol (Life Technologies). The total RNA yield for each sample (average tissue weight of 300 mg) was 200-500 ug. Next, mRNA was isolated using the Oligotex mRNA Midi kit (Qiagen). Since the mRNA was eluted in a final volume of 400 µl, an ethanol precipitation step was required to bring the concentration to 1 µg/µl. Using 1-5 µg of mRNA, double stranded cDNA was created using the SuperScript Choice system (Gibco-BRL). First strand cDNA synthesis was primed with a T7-(dT₂₄) oligonucleotide. The cDNA was then phenol-chloroform extracted and ethanol precipitated to a final concentration of 1 µg/µl.

From 2 µg of cDNA, cRNA was synthesized according to standard procedures. To biotin label the cRNA, nucleotides Bio-11-CTP and Bio-16-UTP (Enzo Diagnostics) were added to the reaction. After a 37°C incubation for six hours, the labeled cRNA was cleaned up according to the Rneasy Mini kit protocol (Qiagen). The cRNA was then fragmented (5× fragmentation buffer: 200 mM Tris-Acetate (pH 8.1), 500 mM KOAc, 150 mM MgOAc) for thirty-five minutes at 94°C.

55 µg of fragmented cRNA was hybridized on the human Hu35k set and the

HuGeneFL array for twenty-four hours at 60 rpm in a 45°C hybridization oven The chips were washed and stained with Streptavidin Phycoerythrin (SAPE) (Molecular Probes) in Affymetrix fluidics stations. To amplify staining, SAPE solution was added twice with an anti-streptavidin biotinylated antibody (Vector Laboratories) staining step in between.

Hybridization to the probe arrays was detected by fluorometric scanning (Hewlett Packard Gene Array Scanner). Following hybridization and scanning, the microarray images were analyzed for quality control, looking for major chip defects or abnormalities in hybridization signal. After all chips passed QC, the data was analyzed using Affymetrix GeneChip software (v3.0), and Experimental Data Mining Tool (EDMT) software (v1.0).

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Example 2: Gene Expression Analysis

All samples were prepared as described and hybridized onto the Affymetrix HuGeneFL array and the Human Hu35k set of arrays. Each chip contains 16-20 oligonucleotide probe pairs per gene or cDNA clone. These probe pairs include perfectly matched sets and mismatched sets, both of which are necessary for the calculation of the average difference. The average difference is a measure of the intensity difference for each probe pair, calculated by subtracting the intensity of the mismatch from the intensity of the perfect match. This takes into consideration variability in hybridization among probe pairs and other hybridization artifacts that could affect the fluorescence intensities. Using the average difference value that has been calculated, the GeneChip software then makes an absolute call for each gene or EST.

The absolute call of present, absent or marginal is used to generate a Gene Signature, a tool used to identify those genes that are commonly present or commonly absent in a given sample set, according to the absolute call. For each set of samples, a median average difference was figured using the average differences of each individual sample within the set. The median average difference must be greater than 150 to assure that the expression level is well above the background noise of the hybridization. For the purposes of this study, only the genes and ESTs with a median average difference greater than 150 have been further studied in detail.

The Gene Signature for one set of samples is compared to the Gene Signature of another set of samples to determine the Gene Signature Differential. This comparison identifies the genes that are consistently present in one set of samples and consistently

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absent in the second set of samples.

The Gene Signature Curve is a graphic view of the number of genes consistently present in a given set of samples as the sample size increases, taking into account the genes commonly expressed among a particular set of samples, and discounting those genes whose expression is variable among those samples. The curve is also indicative of the number of samples necessary to generate an accurate Gene Signature. As the sample number increases, the number of genes common to the sample set decreases. The curve is generated using the positive Gene Signatures of the samples in question, determined by adding one sample at a time to the Gene Signature, beginning with the sample with the smallest number of present genes and adding samples in ascending order. The curve displays the sample size required for the most consistency and the least amount of expression variability from sample to sample. The point where this curve begins to level off represents the minimum number of samples required for the Gene Signature. Graphed on the x-axis is the number of samples in the set, and on the y-axis is the number of genes in the positive Gene Signature.

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Example 3: Gene Expression Analysis of Normal Liver Tissue

The gene expression patterns and Gene Signature were individually determined for each sample set: eight samples with normal liver pathology, six samples whose pathology indicated the primary malignancy to be hepatocellular carcinoma, and five samples whose primary colorectal adenocarcinoma had metastasized to the liver. The Gene Signatures obtained for the normal sample set is shown in Figure 2A, the metastatic liver cancer set in Figure 2B and the hepatocellular carcinoma set in Figure 2C.

The Gene Signature considers the present and absent genes alone, and does not take into consideration those that have been called marginal. Table 2 shows the numbers of present genes, called the positive Gene Signature, and the number of absent genes, called the negative Gene Signature, for each of the three sets of samples.

The Gene Signature is the set of genes that are commonly present or commonly absent in N-1 samples of a given sample set. The positive Gene Signature for the normal liver tissues contains 6,213 genes and ESTs. This same set of normal samples did not show any detectable level of expression of 24,900 genes. Many of the genes and ESTs in this positive Gene Signature are housekeeping genes or structural genes that are not only expressed in the liver, but are ubiquitously expressed in tissues throughout the body. Within

this positive Gene Signature are also those genes whose expression is specifically restricted to normal liver tissue and those genes required for the liver to function at its normal capacities. It is the group of genes unique to the liver whose expression levels are most likely to change during tumorigenesis. Whether up-regulated or down-regulated or turned completely on or turned completely off, the changes in expression of these vital genes very likely contributes to the drastic changes in liver function caused by the transformation of normal liver cells into cancerous cells.

Example 4: Gene Expression Analysis of Malignant Liver Tissue

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There are 8,479 genes and ESTs in the positive Gene Signature for the HCC tumors, and a total of 23,233 genes and ESTs are included in the negative Gene Signature of the HCC samples. This negative Gene Signature includes all the genes that have been completely turned off during tumorigenesis, as well as those genes that are not usually expressed in liver tissue. These results include a number of genes and ESTs that are not regularly expressed in liver tissues, but through the process of tumor production, their expression patterns have been dramatically altered from no detectable level of expression to some significant level of expression in comparison with the normal liver.

The colorectal metastases in the liver commonly express 5,102 genes and ESTs, and do not show expression of 30,455 additional genes and ESTs. As with the negative Gene Signature for the HCC sample set, the genes included in this data set are generally not expressed in liver tissue, whether tumor or normal tissue. The 5,102 in the sample set of metastatic tumors also identify those genes with expression levels that have been changed from off to on as a result of tumor formation.

25 Example 5: Analysis of Gene Expression Profiles

A differential comparison of the genes and ESTs expressed in the normals and the two different types of liver tumors identifies a subset of the genes included in the positive Gene Signatures that are uniquely expressed in each sample set. This Gene Signature Differential highlights genes whose expression profiles have most dramatically changed in the transformation from normal to diseased liver cells. The parameters for these analyses were set to accommodate variation in expression of one of eight normal samples and one of the six HCC samples or one of the 5 metastatic tumor samples, such that the genes

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categorized as unique to normal were called present by the software in seven of eight (87%) normal liver samples and were also called absent in five of six HCC (83%) or four of five (80%) metastatic liver tumor. Conversely, the genes categorized as unique to each set of tumors as compared to the normal livers were called present in five of six HCC (83%) or four of five (80%) metastatic tumor samples and absent in seven of eight normal livers (87%).

The Gene Signature Differential comparing the normal livers to those with metastatic tumors identified a total of 903 sequences expressed only in normal liver tissue. The number of genes or ESTs that meet the median average difference minimum of 150 is 449, of which 289 are genes and the number of ESTs is 160. The remaining ESTs and genes may be indistinguishable from the background noise of the hybridization. The same comparison of normals versus metastatic tumors demonstrates that in the metastatic tumor samples there are 296 uniquely expressed sequences. Those that meet the median average difference minimum requirement are 83 genes and 72 ESTs. Those genes and ESTs expressed in metastatic and not in normal liver tissue are shown in Table 9A and those present in normal liver tissue and not metastatic tissue Table 9B. Numerous genes with differing expression levels in metastatic liver tumor tissue compared to normal tissue were identified. The fifteen genes whose expression level was most different in metastatic as compared to normal tissue are shown in Table 4. Those with the most increased expression are in Table 4A and those with the most decreased expression are in Table 4B. Expression levels were determined by comparing the mean expression values of individual genes in tumor and normal liver samples. Fold change was calculated as a ratio with a p value given as a measure of statistical significance. Fold change is considered significant for a given gene or EST when it is greater than 3.0 with a p value <0.05. Only the characterized genes have been listed; the ESTs with similar fold changes are not presented here. Asterisk (*) in Table 4 denotes those genes that were also identified in the Gene Signature differential between metastatic liver carcinoma and normal liver tissue. A complete listing of all the genes and ESTs with at least a three-fold change in expression is shown in Table 6. Table 6A contains those genes and ESTs whose expression level increased in metastatic tissue relative to normal tissue and Table 6B contains those genes and ESTs whose expression level decreased.

The Gene Signature Differential between the normal liver samples and the HCC

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samples identifies a total of 47 unique expressers in the normals, 23 with an median average difference of 150, 13 of which are named gene and 10 of which are ESTs. When comparing the expression of the HCC samples with the normal livers, there are 243 genes and ESTs only expressed in the HCC samples.

Those genes and ESTs expressed in HCC and not in normal liver tissue are shown in Table 8A and those present in normal liver tissue and not HCC tissue in Table 8B. Numerous genes with differing expression levels in HCC compared to normal tissue were identified. The fifteen genes whose expression level was most different in HCC as compared to normal tissue are shown in Table 3. Those with the most increased expression are in Table 3A and those with the most decreased expression are in Table 3B. Expression levels were determined by comparing the mean expression values of individual genes in tumor and normal liver samples. Fold change was calculated as a ratio with a p value given as a measure of statistical significance. Fold change is considered significant for a given gene or EST when it is greater than 3.0 with a p value <0.05. Only the characterized genes have been listed; the ESTs with similar fold changes are not presented here. Asterisk (*) denotes those genes that were also identified in the Gene Signature differential between hepatocellular carcinoma and normal liver tissue. A complete listing of all the genes and ESTs with at least a three-fold change in expression is shown in Table 7. Table 7A contains those genes and ESTs whose expression level increased in hepatocellular carcinoma tissue relative to normal tissue and Table 7B contains those genes and ESTs whose expression level decreased.

Analysis of sample set identified 24 ESTs and 42 genes that are expressed in both metastatic liver tumors and hepatocellular carcinomas, but not in normal liver tissues. The fifteen genes with the most increase in expression level in both types of cancer are shown in Table 5. Expression levels were determined by comparing the mean expression values of individual genes in tumor and normal liver samples. The mean expression value for HCC and metastatic carcinomas was greater than 250, and included only those genes that showed a fold change greater than 3 with significant p values for both sets of tumors. No detectable level of expression was found in the normal liver tissues for these genes. Only the characterized genes have been listed; the ESTs with similar fold changes that are unique to the tumors are not presented here.

Differential gene expression patterns between normal liver samples and

hepatocellular carcinomas and between normal livers and metastatic liver tumors were examined. Genes uniquely expressed by each of the groups individually were identified, as well as those genes that are commonly expressed among liver tumors, whether primary hepatocellular carcinomas or metastatic liver tumors.

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Example 6: Association of Liver Cancer with Specific Gene Expression

The present inventors have closely examined a number of the tumor-expressing genes to determine if their expression patterns correlate with previous reports published in the literature, and to define a logical relationship between the gene and hepatocarcinogenesis. A number of genes that have previously been associated with either liver cancer or other types of cancers were identified, as well as numerous genes that have not been linked to cancers in any previous studies.

842 genes and ESTs that are up-regulated in hepatocellular carcinomas were identified when compared with normal liver tissue. One such gene is PTTG1, pituitary tumor-transforming gene 1, or securin, an oncogene that inhibits sister chromatid separation during anaphase. Normal tissues show little or no PTTG1 expression, but high levels of expression have been associated with various tumors, including liver tumors, and carcinoma cell lines. Overexpression in NIH3T3 cells resulted in transformation, and these cells caused the formation of tumors when injected into mice. The mechanism by which this tumorigenic activity takes place is postulated to be through the missegregation of sister chromatids, resulting in aneuploidy and, therefore, genetic instability. Our data further support this overexpression of PTTG1 in hepatocellular carcinoma, with a fold change of 10.7 (*P*=0.00052), and no detectable level of expression in normal tissues, as identified by the differential comparison of the consensus patterns of gene expression of these two sample sets.

Galectin 3, LGALS3, one of a family of beta-galactoside-binding animal lectins, is significantly overexpressed both in primary hepatocellular carcinoma and metastatic liver carcinomas with fold changes of 6.8 (P=0.00103) and 27.1 (P=0.00001), respectively. Expression of LGALS3 has been associated with tumor growth, progression, and metastasis, as well as cell-cell and cell-matrix interactions and inflammatory processes. Although expression studies by Hsu *et al.* revealed no detectable level of galectin-3 in normal liver cells, samples from patients with hepatocellular carcinoma revealed considerable levels of

LGALS3 expression. The abnormal expression of this lectin may be an early event in the process of transformation of normal cells to tumor cells, or it may impart an increased capacity for these tumor cells to survive and proliferate. Consistent with the reports by Iurisci et al and Nakamura et al, an increased expression level was found in both types of tumor, but higher concentrations of galectin-3 were observed in liver metastates from colorectal tumors than in the primary HCC tumors.

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Another gene that is overexpressed in both hepatocellular carcinoma and metastatic colorectal adenocarcinomas with fold changes of 12.2 (P=0.00169) and 58.0 (P=0.00063), respectively, is solute carrier family 2, member 3, or glucose transporter 3 (GLUT3). It is one of a family of transmembrane proteins that function as facilitative glucose transporters, which has a unique specificity for brain and neuronal tissues. Glucose uptake and metabolism are known to be increased in carcinoma cells compared to normal cells. Glucose transporter expression may be elevated in response to the increase in glucose utilization seen in actively proliferating cells, like those of tumors. Conversely, the high levels of glucose transporter expression may be responsible for the enhanced influx of glucose into the tumor cells. Various reports have indicated increased expression of one or more of the family of glucose transporters in malignancies, including those of the brain, esophagus, colon, pancreas, liver, breast, lung, bladder, ovary, testis, skin, head and neck, kidney, and gastric tumors. Kurata et al. (Jpn J Cancer Res 1999 Nov;90(11):1238-43) specifically report that metastatic liver carcinomas have even higher levels of GLUT3 expression than primary tumors. Consistent with previous studies, the current data confirm the significant overexpression of GLUT3 both in primary liver cancer, hepatocellular carcinoma, and in tumors that have metastasized from the colon and rectum.

One of the significantly underexpressed genes identified by comparing the expression profiles of hepatocellular carcinomas and metastatic liver tumors with that of normal liver tissue is metallothionein 1L. The expression level in HCC is 26.9 fold lower than that of normal (P=0.00999), and in metastatic colorectal adenocarcinomas it is down-regulated 66.5 fold (P=0.00415). Metallothioneins are heavy metal binding proteins that are involved in detoxification of metals, zinc and copper metabolism cellular adaptation mechanisms, and may be involved in regulating apoptosis. Colorectal adenocarcinoma that has metastasized to the liver has been specifically reported to express less metallothionein than normal liver tissue. Comparison of the consensus patterns of gene expression between

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metastatic liver samples and normal liver samples show no significant level of MT1L expression in the tumors. Furthermore, additional work has determined that human hepatocellular carcinomas contain much lower levels of metallothioneins than normal liver tissue, and that this decrease correlates with the degree of differentiation and concentrations of copper and zinc in the cells. By comparing the expression profiles of hepatocellular carcinoma and normal liver tissue, this significant reduction in MT1L expression in HCC was confirmed.

A number of enzymes belonging to the family of cytochrome P450s are drastically underexpressed in the two sets of liver tumors in comparison with the normal liver tissue. For example, expression of CYP2A6 is decreased in HCC with a fold change of 14.2 (P=0.0307), and in metastatic tumors with a fold change of 69.9 (P=0). CYP8B1 is downregulated 19.3 fold (P=0.00807) in HCC and 65.1 fold (P=0.0039) in liver metastases. In addition to these commonly down-regulated cytochrome P450s, in HCC samples CYP2B is underexpressed 17.9 fold (P=0.01469), and in the metastatic liver tumors CYP2C9 and CYP2A7 are underexpressed 84.7 fold (P=0.00327) and 72.0 fold (P=0), respectively. Several of these genes are also identified by the differential comparison between expression profiles of tumor and normal, confirming the significant decrease in expression in tumor tissues. Many of these P450 enzymes are critical players in the metabolism of carcinogens, drugs, and other chemical compounds, that are expressed in normal liver.

In addition to genes that are underexpressed in metastatic adenocarcinomas in the liver, more than 1000 genes and ESTs that are overexpressed specifically in these tumors were identified. Two of the most highly up-regulated are claudin 4, also known as clostridium perfringens enterotoxin receptor 1 (fold change 84.4, P=0) and occludin (fold change 43.1, P=0). Both of these genes are tight junction proteins, responsible for the formation and maintenance of continuous seals around epithelial cells to form a physical barrier that blocks the free passage of water and solutes through the paracellular space. More specifically, claudin-4 is one member of a family of transmembrane proteins that comprise tight junction strands, and occludin is a cell adhesion molecule (Morita et al., (1999) Proc. Natl. Acad. Sci. U.S.A. 96: 511-516). Claudins likely function as paracellular channels, regulating the flow of ions and solutes into and out of the paracellular space (Simon et al., (1999) Science 235: 103-106; Wong & Goodenough (1999) Science 285: 62; Anderson & Van Itallie (1999) Curr. Biol. 9: R922-924). Tight junction proteins also

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contribute to the regulation of the cellular processes of cell growth and differentiation (Matter & Balda (1999) Int. Rev. Cytol. 186: 117-146). Permeability of tight junctions has been associated with tumor formation, where a breakdown in the barrier function of tight junctions allows an increase in the cellular permeability. This breakdown then opens the tight junction barrier, permitting invasion by tumor cells (Zak et al., (2000) Pflugers Arch. 440: 179-183; Mullin (1997) J. Exp. Zool. 279: 484-489). It has been reported that tight junctions of colon tumors leak more than do the tight junctions of normal colon (Soler et al., (1999) Carcinogenesis 20: 1425-1431). A complete loss of tight junction function and a loss of cell-cell contact growth control was seen in cells that had been transfected with oncogenic Raf-1, and expression levels of occludin and another claudin are lower in these cells (Li & Mrsny (2000) J. Cell Biol. 148: 791-800). Occludin expression has been upregulated in vitro by the addition of various fatty acids that have anti-cancer effects, decreasing the paracellular permeability (Jiang et al., (1998) Biochem. Biophys. Res. Commun. 244: 414-420). The extreme down-regulation of occludin and claudin-4 in metastatic liver tumors is strongly supported by the reports of tight junction breakdown in tumor tissues.

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The present study identified 93 significantly up-regulated genes in both primary HCC and metastatic liver tumors that were not found to have any detectable level of expression in the normal samples. Serine protease inhibitor, Kazal type I (SPINK1), also called pancreatic secretory trypsin inhibitor (PSTI) or tumor-associated trypsin inhibitor (TATI), is one such gene. It is highly expressed in the cells of normal pancreas and in the mucosa of the gastrointestinal tract where it offers protection from proteolytic breakdown. A marked increase in expression is seen in various pancreatic diseases and in tumors of different tissues, including gastric carcinomas, colorectal cancers, and other neoplastic tissues. This increase is presumably due to the elevated expression of trypsin in the tumors, and not related to amplification or rearrangements within the gene. SPINK1 is also considered a valuable marker for a number of solid tumors. A drastic elevation of SPINK1 in the blood of patients with hepatocellular carcinoma has been seen (see Ohmachi et al.). Furthermore, it has been suggested that the level of expression correlates with the extent of tumor, such that this heightened expression level could be indicative of HCC under certain conditions. In keeping with this report of overexpression in these tumors, the present expression data show the levels of expression of this gene in HCC samples to be 28.9 times -33-

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higher than normal (P=0.00003), and in metastatic liver tumors the expression level is 9.8 times higher than normal (P=0.03697).

Midkine is one of a family of heparin-binding growth factors, inducible by retinoic acid, and is actively involved in cell-cell interactions and angiogenesis. The expression pattern of midkine is highly restricted in normal adult tissues, and no expression has been reported in normal adult liver, although its expression is required during embryogenesis for normal development. However, it is expressed in moderate to high levels in many tumors, including Wilm's tumors of the kidney, stomach, colon, pancreas, lung, esophagus, breast, and liver tumors. The present data confirm these reports, showing a significant overexpression of midkine in hepatocellular carcinoma samples (fold change 9.9, P=0.02104) and in liver metastases (fold change 10.4, P=0.01818), but no noticeable expression in normal liver.

Stathmin, leukemia-associated phosphoprotein 18, is a phosphoprotein whose expression pattern and phosphorylation status are controlled by extracellular signals responsible for the regulation of the processes of cell proliferation and differentiation. It is also involved in the regulation of cell division via the destabilization of microtubules. When comparing expression levels between non-malignant tissues and malignant tissues, the tumors generally show a significant up-regulation of this phosphoprotein, specifically lymphomas, leukemias, breast and prostate tumors. One reason proposed for this elevated expression in cancer cells is the dissimilarity in the rates of cell proliferation and states of differentiation between normal and tumor cells. In both HCC samples and metastatic adenocarcinomas, significant up-regulation of stathmin, 9.4 fold in HCC (*P*=0.00015) and 4.8 fold in metastatic tumors (*P*=0.00514) was seen.

Both the genes and ESTs described here will provide valuable information for the identification of new drug targets against liver carcinomas, and that information may be extended for use in the study of carcinogenesis in other tissues.

Although the present invention has been described in detail with reference to examples above, it is understood that various modifications can be made without departing from the spirit of the invention. Accordingly, the invention is limited only by the following claims. All cited patents, applications and publications referred to in this application are herein incorporated by reference in their entirety.

Table 1. Clinical Information for Hepatocellular Carcinoma, Metastatic Liver Tumor, and Normal Liver Samples Included in this Study

| Sample . | Pathology | Primary Tumor | Age | Gender | Race | TNM Classification | Histopathologic Grade | Clinical Stage |
|----------------|---------------------------|---------------|-----|--------|-----------|-----------------------|--------------------------|------------------|
| HCC1 | Hepatocellular Carcinoma | Liver | 64 | Male | Caucasian | T3,Nx,Mx | 62 | stage III |
| HCC2 | Hepatocellular Carcinoma | Liver | 27 | Female | Caucasian | T3,N0,Mx | 55 | stage III |
| 8 | Hepatocellular Carcinoma | Liver | 78 | Female | Caucasian | T4,Nx,Mx | 62 | stage III |
| HCC4 | Hepatocellular Carcinoma | Liver | 43 | Male | Asian | T4,N1,Mx | 62 | stage IV |
| HCC5 | Hepatocellular Carcinoma | Liver | 51 | Male | Caucasian | T4,N0,Mx | G2 | stage (V |
| нссв | Hepatocellular Carcinoma | Liver | 25 | Male | Caucasian | unavailable | 62 | stage III |
| M1 | Metastatic Adenocarcinoma | Rectum | 61 | Female | Caucasian | Tx,Nx,M1 | 63 | stage IV; Duke D |
| M2 | Metastatic Adenocarcinoma | Colon | 54 | Male | Caucasian | unavailable | 62 | stage IV; Duke D |
| M3 | Metastatic Adenocarcinoma | Rectum | 20 | Female | Caucasian | Tx,Nx,M1 | 62 | stage IV; Duke D |
| M4 | Metastatic Adenocarcinoma | Colon | 09 | Maie | Caucasian | Tx,Nx,M1 | G 2 | stage IV; Duke D |
| M5 | Metastatic Adenocarcinoma | Colon | 25 | Male | Caucasian | Tx,Nx,M1 | 62 | stage IV; Duke D |
| Ξ | Normal liver | | 54 | Female | Caucaslan | | | |
| ~ | Normal liver | | 22 | Female | Caucasian | | | |
| 5 | Normal liver | | 28 | Male | Caucasian | - | | |
| * | Normal liver | | 44 | Female | Caucasian | | | |
| 55 | Normal liver | - | 40 | Female | Caucasian | | | |
| Ne Ne | Normal liver | | 72 | Female | Caucasian | | | |
| N ₂ | Normal liver | | 48 | Female | Unknown | | | |
| 88 88 | Normal liver | | 22 | Female | Caucasian | | | |

Table 2. Summary of Genes and ESTs Expressed in HCC, Metastatic Liver Tumors, and Normal Livers

| | Hepatocellular Carcinoma | Colorectal Metastases | Normal Livers |
|---|-----------------------------|--------------------------|---------------|
| I. Fingerprint of Gene Expression | | | |
| Genes and ESTs commonly expressed in sample set | t 8479 | 5102 | 6213 |
| Genes and ESTs commonly unexpressed in sample set | t 23233 | 30455 | 24900 |
| II. Fold Change | | | |
| Genes and ESTs overexpressed in tumors (fold change >3 and p<0.05) | 842 | 1044 | |
| Number of Genes | 430 | 603 | |
| Number of ESTs | 3 412 | 441 | |
| Genes and ESTs underexpressed in tumors (fold change >3 and p<0.05) | . 393 | 1867 | |
| Number of Genes | 235 | 1016 | |
| Number of ESTs | 158 | 851 | |
| | | | |
| III. Differential Comparison between Normal and Tumor Expression | | | |
| Genes and ESTs turned ON in turnors | ••• | 296 | |
| Genes and ESTs with expression level above threshold in tumor | r 77 | 155 | |
| Number of Genes | | 83 | |
| Number of ESTs | 39 | 72 | |
| Genes and ESTs turned OFF in turnors | 3 47 | 903 | |
| Genes and ESTs with expression level above threshold in normal | 23 | 449 | |
| . Number of Genes | 13 | 289 | |
| Number of ESTs | 10 | 160 | |

Table 3A. Top fifteen genes overexpressed in hepatocellular carcinoma.

| Genbank | Seq ID | Unigene Cluster | ter Gene Name | Fold Change Pyalue | Pyalma |
|----------|--------|-----------------|---|--------------------|----------|
| AA055896 | 135 | Hs.146428 | collagen, type V, alpha 1 | 10.9* | 0 00907 |
| AA156187 | 339 | Hs.81634 | ATP synthase, H+ transporting, mitochondrial F0 complex, subunit b, isoform 1 | 76 | |
| AA428172 | 986 | Hs.8546 | Notch (Drosophila) homolog 3 | *90 | |
| AA430032 | 1009 | Hs.252587 | pituitary tumor-transforming 1 | 10.7* | 0.00183 |
| AA505133 | 1417 | Hs.279905 | solute carrier family 2 (facilitated glucose transporter), member 3 | 12.2 | _ |
| AA610116 | .1499 | Hs.102737 | tetraspan NET-6 protein | 16.4 | 0.00103 |
| AA620881 | 1510 | Hs.21858 | trinucleotide repeat containing 3 | 1.0 | 0.0062 |
| D31094 | 1639 | Hs.109798 | G8 protein | 5.0 | • |
| D51276 | 1678 | Hs.81915 | leukemia-associated phosphoprotein p18 (stathmin) | t 5 | C |
| J03464 | 2094 | Hs.179573 | collagen, type I, alpha 2 | 7 0.1 | |
| M94250 | 2426 | Hs.82045 | midkine (neurite arowth-promotina factor 2) | *00 | 0.0037.9 |
| N33920 | 2493 | Hs.44532 | diubiauitin | 503 | |
| W45320 | 3523 | Hs.228059 | KRAB-associated protein 1 | 10 1* | |
| Y00705 | 3850 | Hs.181286 | serine profease inhibitor. Kazal tyne 1 | 280 | , , |
| Z37987 | 3882 | Hs.119651 | glypican 3 | 10.7 | 0.02304 |

Table 3B. Top fifteen genes underexpressed in hepatocellular carcinoma.

| Genbank | Seq ID | Unigene Cluster | Gene Name | Fold Change Pvalue | Pvalue |
|----------|--------|-----------------|---|--------------------|--------------|
| AA007395 | 17 | Hs.1219 | alcohol dehydrogenase 4 (class II), pi polypeptide | 37.8 | 37.8 0.00939 |
| AA010605 | 26 | Hs.2899 | 4-hydroxyphenylpyruvate dioxygenase | 25.5 | 25.5 0.00855 |
| AA448002 | 1113 | Hs.23759 | putative type II membrane protein | 14.1* | 0 |
| H58692 | 1960 | Hs.9520 | formyltetrahydrofolate dehydrogenase | 20.2 | 20.2 0.00485 |
| H80901 | 2005 | Hs.272576 | ficolin (collagen/fibrinogen domain-containing) 3 (Hakata antigen) | 18.6 | 0 |
| H81070 | 2006 | Hs.8765 | RNA helicase-related protein | 39.6 | 39.6 0.00002 |
| K03192 | 2127 | Hs/183584 | cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 | 14.2 | 0.0307 |
| M29873 | 2318 | Hs.1360 | cytochrome P450, subfamily IIB (phenobarbital-inducible) | 17.9 | 0.01469 |
| N80129 | 2703 | Hs.94360 | metallothionein 1L | 26.9 | 0.00999 |
| R97419 | 3004 | Hs.35718 | cytochrome P450, subfamily VIIIB (sterol 12-alpha-hydroxylase), polypeptide 1 | 19.3 | 0.00807 |
| T48075 | 3130 | Hs.251577 | hemoglobin, alpha 1 | 35.8 | 0.00471 |
| T67931 | 3184 | Hs.7645 | fibrinogen, B beta polypeptide | 17.3 | 0.00128 |
| T95813 | 3262 | Hs.137476 | KIAA1051 protein | 20.4 | 0.01361 |
| U56814 | 3393 | Hs.88646 | deoxyribonuclease I-like 3 | 17.7 | 0.00007 |
| W88946 | 3639 | Hs.18508 | putative glycine-N-acyltransferase | 25.3 | 25.3 0.00221 |

Table 4A. Top fifteen genes overexpressed in metastatic carcinomas of the liver.

| Genbank | Seq ID | Unigene Cluster | Gene Name | Fold Change Dyalna | Pyshia |
|---------------|--------|-----------------|---|--------------------|--------------|
| AA100719 | 212 | Hs.73848 | Carcinoembryonic antigen-related cell adhesion molecule 6 (non-specific | 50.9* | 0.00081 |
| | | | cross reacting antigen) | | |
| AA156243 | 340 | Hs.154737 | Serine protease, umbilical endothelium | 41.4* | 0.00139 |
| AA335191 | 741 | Hs.173724 | Creatine kinase, brain | 47.4* | 0.00449 |
| AA421562 | 934 | Hs.91011 | Anterior gradient 2 (Xenopus leavis) homolog | 56.3* | 0.001 |
| AA427468 | 973 | Hs.5372 | Claudin 4 | 84.4* | - C |
| AA429009 | 994 | Hs.233950 | Serine protease inhibitor, Kunitz type 1 | 30.0* | 0 00001 |
| AA610116 | 1499 | Hs.102737 | Tetraspan NET-6 protein | | 33.7 0.00171 |
| H58873 | 1961 | Hs.169902 | Solute carrier family 2 (facilitated glucose transporter), member 3 | 58.0* | 0.00063 |
| H94471 | 2042 | Hs.171952 | Occludin | 73.1 | |
| H95233 | 2048 | Hs.31439 | Serine protease inhibitor. Kunitz tyne 2 | 17.7 | · • |
| HG2788-HT2896 | | | Calcyclin binding protein | 33.0 | - ^ |
| M29540 | 2317 | 6 | Carcinoembryonic antigen-related cell adhesion molecule 5 | * 99 | 5 0 0118 |
| M35252 | 2343 | Hs.84072 | Transmembrane 4 superfamily member 3 | 39.1 | |
| N92934 | 2725 | Hs.17409 | Cysteine-rich protein 1 (intestinal) | 25. 25.5 | |
| X93036 | 3830 | Hs.92323 | FXYD domain-containing ion transport regulator 3 | 42.4* | Ö |

Table 4B. Top fifteen genes underexpressed in metastatic carcinoma of the liver.

| Genbank | Seq ID | Unigene Cluste | ter Gene Name | Fold Change Pvalue | Pvalue |
|----------|--------|----------------|---|--------------------|--------------|
| AA256367 | 579 | Hs.107966 | Paraoxonase 3 | 3.07 | 70.3 0.00192 |
| H58692 | 1960 | Hs.9520 | Formyltetrahydrofolate dehydrogenase | 81.4* | 0 |
| K03192 | 2127 | Hs.183584 | Cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 | *6.69 | 0 |
| L16883 | 2166 | Hs.167529 | Cytochrome P450, subfamily IIC (mephenytoin 4-hydroxylase), polypeptide 9 | 84.7 | 84.7 0.00327 |
| M15656 | 2268 | Hs.234234 | Aldolase B, fructose-bisphosphate | 36.7 * | 0 |
| M16594 | 2272 | Hs.89552 | glutathione S-transferase A2 | 73.2* | 0 |
| M81349 | 2405 | Hs.1955 | Serum amyloid A4, constitutive | 76.2 | 76.2 0.00015 |
| N53031 | 2556 | Hs.89691 | UDP glycosyltransferase 2 family, polypeptide B4 | 9.76 | 3 0.00022 |
| N54417 | 2567 | Hs.90765 | Fibrinogen, A alpha polypeptide | 99.3 | 3 0.00001 |
| R43174 | 2848 | Hs.1898 | Paraoxonase 1 | 74.0* | 0.00038 |
| R49459 | 2882 | Hs.63758 | Transferrin receptor 2 | 85.6 | 3 0.00048 |
| T48039 | 3129 | Hs.2351 | Protein C (inactivator of coagulation factors Va and VIIIa) | 84.4 | 1 0.00112 |
| T59148 | 3158 | Hs.50966 | Carbamoyl-phosphate synthetase 1, mitochondrial | 88.9* | 0 |
| U22029 | 3327 | Hs.250615 | Cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 7 | 72.0* | 0 |
| X55283 | 3734 | Hs.1259 | Asialoglycoprotein receptor 2 | 86 | 85 0.00084 |

Table 5. Top fifteen genes expressed in both hepatocellular carcinomas and metastatic liver tumors ad not in normal livers.

| Genbank | Seq ID | Unigene Cluster | Gene Name | HCC Fold | HCC p | Metastatics Metastatic | Metastatic |
|----------|--------|-----------------|---|----------|---------|------------------------|------------|
| | | | | Change | value | Fold | p value |
| | | | | | | Change | |
| AA055896 | 135 | Hs.146428 | collagen, type V, alpha 1 | 10.9 | 0.00907 | 18.2 | 0.00146 |
| AA204927 | 425 | Hs.77899 | tropomyosin 1 (alpha) | 6.1 | 0.0014 | 7.1 | 0.00074 |
| AA335191 | 741 | Hs.173724 | creatine kinase, brain | 6.5 | 0.01462 | 47.4 | 0.00419 |
| AA429472 | 266 | Hs.236522 | DKFZP434P106 protein | 8.8 | 0.00063 | 80 | 0.00208 |
| AA434418 | 1036 | Hs.72172 | KIAA1115 protein | 6.8 | 0.0032 | 5.1 | 0.00498 |
| AA452724 | 1149 | Hs.166468 | programmed cell death 5 | 7.7 | 0.00085 | 7.2 | 0.00908 |
| AA610116 | 1499 | Hs.102737 | tetraspan NET-6 protein | 16.4 | 0.00249 | 33.7 | 0.00171 |
| AA620881 | 1510 | Hs.21858 | trinucleotide repeat containing 3 | 9.5 | 0.00062 | 8.7 | 0.00735 |
| D26129 | 1635 | Hs.78224 | ribonuclease, RNase A family, 1 (pancreatic) | 6.9 | 0.00008 | 5.7 | 0.03827 |
| D31094 | 1639 | Hs.109798 | G8 protein | 9.4 | 0.0048 | 4.4 | 0.04845 |
| D51276 | 1678 | Hs.81915 | leukemia-associated phosphoprotein p18 (stathmin) | 9.4 | 0.00015 | 8.4 | 0.00514 |
| H27188 | 1908 | Hs.9930 | collagen-binding protein 2 (colligen 2) | 5.8 | 0.01826 | 4.2 | 0.02073 |
| J03464 | 2094 | Hs.179573 | collagen, type I, alpha 2 | 10.4 | 0.00979 | 8 | 0.00028 |
| M94250 | 2426 | Hs.82045 | midkine (neurite growth-promoting factor 2) | 6.6 | 0.02104 | 10.4 | 0.01818 |
| Y00705 | 3850 | Hs.181286 | serine protease inhibitor, Kazal type 1 | 28.9 | 0.00003 | 80 | 0.03697 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|--------------|---------|
| rc_AA001409_i_at | AA001409 | - | EST | 3.35 | dn | 0.04092 |
| rc_AA001504_f_at | AA001504 | 7 | EST | 9.98 | dn | 0.00336 |
| rc_AA007158_f_at | AA007158 | 15 | EST | 3.05 | dn | 0.01964 |
| rc_AA007160_at | AA007160 | 16 | EST | 9 | dn | 0.01035 |
| rc_AA010065_s_at | AA010065 | 22 | CDC28 protein kinase 2 | 6.25 | ď | 0.02752 |
| rc_AA011134_at | AA011134 | 59 | EST | 28.79 | d | 0.00602 |
| rc_AA011383_at | AA011383 | 31 | EST | 5.17 | 요 | 0.00008 |
| rc_AA025166_s_at | AA025166 | 20 | fusion, derived from t(12;16) malignant liposarcoma | 3.71 | dn . | 0.0052 |
| rc_AA025277_at | AA025277 | 5. | EST | 4.56 | dn | 0.03136 |
| rc_AA026030_at | AA026030 | 23 | EST | 11.01 | dn | 0.01649 |
| rc_AA026092_at | AA026092 | 54 | EST | 3.83 | dn | 0.04596 |
| rc_AA026150_at | AA026150 | 22 | EST | 5.14 | ď | 0.01072 |
| rc_AA026356_at | AA026356 | 22 | EST | 4.1 | dn | 0.00133 |
| rc_AA027946_at | AA027946 | 09 | EST | 3.22 | dn | 0.00098 |
| rc_AA028103_at | AA028103 | 61 | EST | 3.52 | g | 0.01142 |
| rc_AA028132_s_at | AA028132 | 62 | EST | 6.25 | 슠 | 0.00646 |
| rc_AA029215_at | AA029215 | .04 | adaptor-related protein complex 2, beta 1 subunit | 3.65 | dn | 0.00037 |
| rc_AA029356_at | AA029356 | 99 | EST | 3.68 | dn | 0.01545 |
| rc_AA033790_f_at | AA033790 | 74 | apolipoprotein D | 4.21 | <u>a</u> | 0.03247 |
| rc_AA034378_f_at | AA034378 | 11 | endogenous retroviral protease | 4.01 | dn | 0.00974 |
| rc_AA034499_s_at | AA034499 | 78 | zinc finger protein 198 | 3.7 | dn | 0.02143 |
| rc_AA040465_at | AA040465 | 92 | EST | 3.25 | dn | 0.00146 |
| rc_AA043959_at | AA043959 | 101 | tropomyosin 4 | 8.54 | dn | 0.00222 |
| rc_AA053007_f_at | AA053007 | 123 | putative receptor protein | 9.76 | dn | 0.00061 |
| rc_AA053033_at | AA053033 | 124 | EST | 7.83 | dn | 0.00379 |
| rc_AA053102_s_at | AA053102 | 125 | cadherin 17, Ll cadherin (liver-intestine) | 26.63 | dn | 0.01745 |
| rc_AA053248_l_at | AA053248 | 126 | EST | 7.01 | dn | 0.00003 |
| rc_AA053248_f_at | AA053248 | 126 | EST | 6.16 | dn | 0.00191 |
| rc_AA053424_at | AA053424 | 127 | EST | 8.76 | dn | 0.01775 |
| rc_AA053660_at | AA053660 | 128 | EST | 15.98 | dn | 0.00003 |
| rc_AA055805_s_at | AA055805 | 132 | EST | 42.83 | dn | 0.00142 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| rc_AA055811_s_at | AA055811 | 133 | glycoprotein A33 (transmembrane) | 98.9 | dn | 0.02152 |
| rc_AA055896_at | AA055896 | 135 | collagen, type V, alpha 1 | 18.16 | dn | 0.00146 |
| AA056361_at | AA056361 | 140 | integral membrane protein 2C | 3.53 | dn | 0.02983 |
| rc_AA062721_at | AA062721 | 146 | nuclear factor (erythroid-derived 2)-like 1 | 6.2 | dn | 0.00024 |
| rc_AA070206_at | AA070206 | 155 | EST | 4.26 | dn | 0.00018 |
| rc_AA070827_at | AA070827 | 157 | EST | 4.41 | dn | 0.01902 |
| rc_AA074514_at | AA074514 | 160 | EST | 7.69 | dn | 0 |
| rc_AA075299_at | AA075299 | 164 | EST | 34.27 | dn | 0.00002 |
| rc_AA075580_f_at | AA075580 | 165 | EST | 4.98 | dn | 0.02083 |
| rc_AA075722_at | AA075722 | 166 | nuclear transport factor 2 (placental protein 15) | 4.14 | dn | 0.00374 |
| AA078862 s at | AA078862 | 173 | EST | 6.52 | dn | 0.00025 |
| rc_AA084901_at | AA084901 | 181 | ribosomal protein S6 kinase, 70kD, polypeptide 2 | 9.13 | dn | 0.00003 |
| rc_AA084921_f_at | AA084921 | 182 | ribosomal protein S10 | 3.29 | dn | 0.04872 |
| | | | kangai 1 (suppression of tumorigenicity 6, prostate; CD82 | | | |
| rc AA086232 f at | AA086232 | 186 | and antihody [A41] | 5 14 | 9 | 0.01916 |
| AA093497 s at | AA093497 | 199 | DEK oncodene (DNA bindina) | 5.5 | t 8 | 0.02551 |
| AA094517 at | AA094517 | 202 | EST | 3.52 | ٠ 1 | 0.04805 |
| AA094752 at | AA094752 | 203 | hypothetical 43.2 Kd protein | 4.47 | d a | 0.00243 |
| rc_AA099404 s_at | AA099404 | 208 | EST | 29.07 | G | 0 |
| i } | | | carcinoembryonic antigen-related cell adhesion molecule 6 (non- | | • | |
| rc_AA100719_s_at | AA100719 | 212 | specific cross reacting antigen) | 50.88 | dn | 0.00081 |
| rc_AA101551_at | AA101551 | 216 | EST | 24.78 | dn | 0 |
| rc_AA113149_s_at | AA113149 | 226 | tumor suppressing subtransferable candidate 3 | 10.58 | dn | 0.00543 |
| rc_AA113303_at | AA113303 | 227 | transmembrane 4 superfamily member (tetraspan NET-7) | 3.73 | dn | 0.00084 |
| rc_AA116036_at | AA116036 | 233 | chromosome 20 open reading frame 1 | 8.81 | dn | 0.00133 |
| rc_AA121315_at | AA121315 | 237 | KIAA1077 protein | 4.86 | dn | 0.02438 |
| rc_AA122386_at | AA122386 | 239 | collagen, type V, alpha 2 | 4.28 | dn | 0.005 |
| rc_AA126044_at | AA126044 | 245 | EST | 11.23 | dn | 0.00041 |
| rc_AA126429_at | AA126429 | 247 | peroxisomal farnesylated protein | 3.23 | ф | 0.00478 |
| | | | | | | |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| DKFZP566B023 protein 3.69 up EST 6.59 up EST 3.23 up EST 3.23 up EST 3.78 up procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3 4.74 up procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3 4.74 up EST 6.09 up DKFZP564O0463 protein 3.76 up EST 3.76 up DKFZP564O0463 protein 4.19 up EST 3.31 up EST 9.72 up EST 9.19 up EST 9.19 up EST 12.11 up endogenous retroviral protease 3.64 up EST 12.11 up pleckstrin 2 (mouse) homolog | Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | le Direction | Pvalue |
|--|------------------|----------|--------|--|-------------|--------------|---------|
| AA/128719 256 EST AA/12871 5.56 Up AA/128712 255 EST 6.59 Up AA/12861 267 EST 3.78 Up AA/12861 266 EST 3.78 Up AA/131684 266 EST 4.74 Up AA/131694 266 EST 4.74 Up AA/131694 268 EST 4.74 Up AA/131694 269 EST 4.74 Up AA/131694 276 DKFZP564O0463 protein 6.63 Up AA/131694 276 DKFZP564O0463 protein 6.63 Up AA/131694 276 DKFZP564O0463 protein 3.76 Up AA/131694 276 DKFZP566G1517 protein 3.76 Up AA/132696 277 EST AA/13269 AA/13269 Up AA/132696 278 EST AA/13269 AA/13269 AA/13269 Up AA/132696 | | AA126459 | 248 | | 3.69 | ı — | 0.00352 |
| att AA127851 255 EST 6.59 up att AA128651 267 EST 3.23 up att AA128651 267 EST 3.78 up att AA131084 265 procollegen-lysine, 2-oxoglutarate 5-dioxygenase 3 4.74 up att AA131084 266 EST 4.74 up att AA131084 266 EST 4.74 up att AA131094 269 EST 4.74 up att AA131094 269 EST 4.74 up att AA131094 270 putative type II membrane protein 6.83 up att AA132032 271 trinuclostide repeat containing 1 4.19 up att AA132032 273 EST 4.96 up att AA132083 274 DKFZP586G1517 protein 3.37 up att AA133080 285 EST 4.10 up s_att AA134065 286 EST 4.26 up att AA134065 286< | rc_AA126719_at | AA126719 | 250 | EST | 5.31 | <u>d</u> | 0.00026 |
| att AA/12861 257 EST att AA/12861 259 EST att AA/12861 265 procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3 3.78 up s_att AA/131684 265 procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3 4.74 up att AA/131684 266 EST 4.74 up att AA/131684 266 EST 4.74 up att AA/131684 266 EST 4.74 up s_att AA/131694 270 putative type II membrane protein 4.78 up s_att AA/132654 273 EST 4.19 up att AA/132654 273 EST 4.19 up att AA/132664 275 EST 4.19 up att AA/132664 275 EST 4.19 up att AA/132664 273 EST 4.19 up att | AA127712_at | AA127712 | 255 | EST | 6.59 | dn | 0.03706 |
| ## AA128407 259 EST ## AA128407 259 EST ## AA131684 266 Forcollagen-tysine, 2-oxoglutarate 5-dioxygenase 3 | - 1 | AA127851 | 257 | EST | 3.23 | dn | 0.01943 |
| att AA/128561 261 collagen, type XVII, alpha 1 6.09 up s_att AA/13162 266 EST cocllagen-lysine, 2-oxoglutarate 5-dioxygenase 3 4.74 up s_att AA/13162 268 EST procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3 4.74 up att AA/13164 268 EST 4.68 up att AA/131894 269 EST up att AA/131894 269 EST up att AA/13264 273 EST up att AA/13289 274 DKFZP586C1517 protein 4.19 up att AA/13289 275 EST 4.99 up att AA/13289 275 EST 4.19 up att AA/13289 285 Rab geranylgeranyltransferase, alpha subunit 3.54 up att AA/13468 287 EST 12.11 up att AA/13468 287 EST 282 | | AA128407 | 259 | EST | 3.78 | dn | 0.01081 |
| at AA(131084 265 procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3 4.74 up sat AA(13162 266 EST 4.68 up at AA(131694) 268 DKFZP564O0463 protein 6.83 up at AA(131694) 270 putative type II membrane protein 4.68 up at AA(132695 277 trinucleotide repeat containing 1 4.19 up at AA(132986 277 EST 4.96 up at AA(132986 275 EST 4.96 up at AA(132986 275 EST 4.96 up at AA(132986 275 EST 4.96 up at AA(13080 285 Rab geran/geran/granyftransferase, alpha subunit 3.23 up s_at AA(134086 289 EST EST up at AA(134086 289 EST EST up at AA(135694 296 | rc_AA128561_at | AA128561 | 261 | collagen, type XVII, alpha 1 | 60.9 | dn | 0.0433 |
| s_att AA131162 266 EST 4.68 up att AA131844 268 DKFZP66400463 protein 6.83 up att AA131894 269 EST up att AA132032 271 trinucleotide repeat containing 1 4.19 up att AA132032 277 trinucleotide repeat containing 1 4.19 up att AA132032 277 trinucleotide repeat containing 1 4.19 up att AA132036 275 EST 4.19 up att AA132096 275 EST up 4.19 up att AA13209 282 EST up 9.72 up att AA13309 284 EST up 9.72 up att AA13405 285 Rab geran/geran/gransferase, alpha subunit 3.54 up att AA13406 286 EST andogenous retroviral protease 28 up | rc_AA131084_at | AA131084 | 265 | procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3 | 4.74 | dn | 0.00427 |
| at AA131584 268 DKFZP56400463 protein 6.83 up at AA131894 269 EST ap at AA132919 270 putative type II membrane protein 15.36 up at AA132963 271 trinucleotide repeat containing 1 4.19 up at AA132983 274 DKFZP586G1517 protein 4.96 up at AA132986 275 EST up 4.96 up at AA132986 275 EST up 4.72 up at AA132986 275 EST up 9.72 up at AA134082 284 EST ap 9.72 up at AA134062 285 EST ap 9.72 up at AA134062 286 EST ap 9.72 up at AA134062 286 EST ap 12.11 up at <th< td=""><td>rc_AA131162_s_at</td><td>AA131162</td><td>266</td><td>EST</td><td>4.68</td><td>dn</td><td>0.00042</td></th<> | rc_AA131162_s_at | AA131162 | 266 | EST | 4.68 | dn | 0.00042 |
| at AA131894 269 EST 3.76 up Leat AA132092 271 trinucleotide repeat containing 1 AA132093 274 breezest containing 1 AA132096 275 EST AA132090 282 EST AA133090 282 EST AA133090 282 EST AA133090 282 EST AA134062 289 EST AA134062 289 EST AA134068 289 EST AA134068 289 EST AA134068 289 EST AA134068 289 EST AA134070 292 endogenous retroviral protease at AA13407 292 endogenous retroviral protease AA136470 302 EST AA136470 302 EST AA143763 310 EST AA143763 311 EST AA143763 312 EST AA143763 312 EST AA143764 315 EST AA147084 BST AA147084 315 EST AA147084 316 AA147084 316 BST AA147084 316 AA147084 3 | rc_AA131584_at | AA131584 | 268 | DKFZP56400463 protein | 6.83 | dn | 0.00025 |
| at AA131919 270 putative type II membrane protein 15.36 up s_at AA132032 271 trinucleotide repeat containing 1 4.19 up at AA132983 274 DKFZP586G1517 protein 4.96 up at AA132983 275 EST AA13399 282 Up at AA13399 282 EST Up 9.72 up at AA13399 284 EST Up 9.19 up s_at AA134052 285 Rab geranylgeranyltransferase, alpha subunit 3.54 up s_at AA134052 286 Rab geranylgeranyltransferase, alpha subunit 3.54 up s_at AA134052 286 Rab geranylgeranyltransferase, alpha subunit 3.54 up s_at AA134062 286 EST up 3.24 up at AA134068 289 EST Up 3.54 up at AA136647 296 EST | rc_AA131894_at | AA131894 | 269 | EST | 3.76 | dn | 0.00384 |
| s_att AA132032 271 trinucleotide repeat containing 1 4.19 up at AA132554 273 EST 4.96 up at AA132983 274 DKFZP586G1517 protein 4.96 up at AA13393 284 EST up at AA13359 285 EST up s_at AA134062 286 Rab geranylgeranyltransferase, alpha subunit 3.54 up s_at AA134062 286 Rab geranylgeranyltransferase, alpha subunit 3.54 up at AA134068 289 EST up 12.11 up at AA134068 289 EST up 12.11 up at AA13681 296 EST and and and at AA13684 296 retinoic acid induced 3 acid acid acid at AA13684 296 retinoic acid induced 3 acid acid acid | rc_AA131919_at | AA131919 | 270 | putative type II membrane protein | 15.36 | dn | 0.00027 |
| at AA132554 273 EST 4.96 up at AA132883 274 DKFZP586G1517 protein 3.31 up at AA1332886 275 EST up at AA133590 282 EST up at AA133596 284 EST up at AA134052 285 Rab geranylgeranyltransferase, alpha subunit 3.54 up at AA134062 285 EST up at AA134068 289 EST up at AA134088 299 EST up at AA135871 294 EST 12.11 up at AA135894 295 retinoic acid induced 3 3.64 up at AA135894 295 retinoic acid induced 3 3.54 up at AA135894 295 retinoic acid induced 3 3.51 up at AA135894 295 retinoic acid induced 3 | rc_AA132032_s_at | AA132032 | 271 | trinucleotide repeat containing 1 | 4.19 | dn | 0.00295 |
| at AA132983 274 DKFZP586G1517 protein 3.31 up at AA132986 275 EST up at AA133590 282 EST up at AA134052 284 EST up as_at AA134052 285 Rab geranylgeranyltransferase, alpha subunit 3.54 up at AA134052 285 Rab geranylgeranyltransferase, alpha subunit 3.54 up at AA134052 285 Rab geranylgeranyltransferase, alpha subunit 3.54 up at AA134085 290 EST up at AA135871 292 endogenous retroviral protease 3.64 up at AA135871 294 EST 3.54 up at AA135871 294 EST 3.54 up at AA135894 295 retinoic acid induced 3 3.51 up at AA143493 310 pleckstrin 2 (mouse) homolog 4.58 < | rc_AA132554_at | AA132554 | 273 | EST | 4.96 | dn | 0.02076 |
| at AA132986 275 EST up at AA133590 282 EST 3.23 up at AA133590 284 EST 9.19 up s_at AA134052 285 Rab geranylgeranyltransferase, alpha subunit 3.54 up s_at AA134052 285 Rab geranylgeranyltransferase, alpha subunit 3.54 up at AA134058 289 EST 12.11 up at AA134085 290 EST 12.11 up AA135871 294 EST 3.34 up at AA135894 295 retinoic acid induced 3 3.39 up at AA135894 295 retinoic acid induced 3 3.51 up at AA143693 310 pieckstrin 2 (mouse) homolog 4.58 up at AA1476619 312 EST 4.4 up at AA147084 314 proliferation-associated 2G4, 38kD 4.4 </td <td>rc_AA132983_at</td> <td>AA132983</td> <td>274</td> <td>DKFZP586G1517 protein</td> <td>3.31</td> <td>dn</td> <td>0.01155</td> | rc_AA132983_at | AA132983 | 274 | DKFZP586G1517 protein | 3.31 | dn | 0.01155 |
| at AA133590 282 EST at AA133590 282 EST at AA13936 284 EST 9.19 up s_at AA134052 285 Rab geranylgeranyltransferase, alpha subunit 3.54 up s_at AA134968 287 EST up at AA134985 290 EST 12.11 up f_at AA135871 294 EST 14.27 up at AA135871 294 EST 3.39 up at AA135877 295 retinoic acid induced 3 3.39 up at AA135894 295 retinoic acid induced 3 3.51 up at AA135877 302 EST at AA13589 4.58 up at AA136547 302 EST at AA146619 312 EST up at AA146619 314 proliferation-associated 2G4, 38kD 4.4 up | rc_AA132986_at | AA132986 | 275 | EST | 9.72 | dn | 0.00552 |
| at AA133936 284 EST 9.19 up s_at AA134052 285 Rab geranylgeranyltransferase, alpha subunit 3.54 up s_at AA13405 287 EST up at AA134968 289 EST 12.11 up f_at AA135871 292 endogenous retroviral protease 3.64 up at AA135871 294 EST up at AA135894 295 retinoic acid induced 3 3.39 up at AA135847 302 EST up 4.58 up at AA143657 302 EST at 4.58 up at AA143763 311 EST 4.58 up at AA147684 312 EST 4.4 up at AA1477084 314 proliferation-associated 2G4, 38kD 4.4 up s_at AA1477439 315 EST 4.4 up< | rc_AA133590_at | AA133590 | 282 | EST | 3.23 | dn | 0.03565 |
| s_at AA134052 285 Rab geranylgeranyltransferase, alpha subunit 3.54 up at AA134158 287 EST up at AA134985 290 EST 12.11 up f_at AA135407 292 endogenous retroviral protease 3.64 up at AA135871 294 EST up 3.39 up at AA135894 295 retinoic acid induced 3 3.51 up at AA136547 302 EST 4.58 up at AA143693 310 pleckstrin 2 (mouse) homolog 4.58 up at AA143763 311 EST 4.87 up at AA147084 312 EST 4.87 up s_at AA147439 315 EST A.97 up s_at AA147439 315 EST A.97 A.97 | rc_AA133936_at | AA133936 | 284 | EST | 9.19 | dn | 0.00088 |
| s_at AA134158 287 EST up at AA134968 289 EST 12.11 up fat AA134985 290 EST 12.11 up fat AA135407 292 endogenous retroviral protease 3.64 up at AA135894 295 retinoic acid induced 3 3.39 up at AA136547 302 EST 4.27 up at AA143493 310 pleckstrin 2 (mouse) homolog 4.58 up at AA143763 311 EST 18.09 up at AA146619 312 EST 4.4 up at AA147084 314 proliferation-associated 2G4, 38kD 4.4 up s_at AA147084 315 EST 5.01 up | | AA134052 | 285 | Rab geranylgeranyltransferase, alpha subunit | 3.54 | dn | 0.03062 |
| at AA134968 289 EST 12.11 up _at AA134985 290 EST 12.11 up _at AA135407 292 endogenous retroviral protease 3.64 up _at AA135871 294 EST 3.39 up _at AA135894 295 retinoic acid induced 3 14.27 up _at AA136547 302 EST 3.51 up _at AA143493 310 pleckstrin 2 (mouse) homolog 4.58 up _at AA143763 311 EST 18.09 up _at AA146619 312 EST 4.4 up _s_at AA147084 314 proliferation-associated 2G4, 38kD 4.4 up _s_at AA147439 315 EST 5.01 up | ٠٠, | AA134158 | 287 | EST | 3.42 | dn | 0.0277 |
| at AA134985 290 EST f_at AA135407 292 endogenous retroviral protease 3.64 up at AA135871 294 EST 3.39 up at AA136547 302 EST 4.27 up at AA143493 310 pleckstrin 2 (mouse) homolog 4.58 up at AA143763 311 EST 4.87 up at AA146619 312 EST 4.4 up s_at AA147084 314 proliferation-associated 2G4, 38kD 4.4 up s_at AA147439 315 EST 5.01 up | rc_AA134968_at | AA134968 | 289 | EST | 12.11 | dn | 0.00079 |
| f at AA135407 292 endogenous retroviral protease 3.64 up at AA135871 294 EST 3.39 up at AA135894 295 retinoic acid induced 3 14.27 up at AA136547 302 EST 3.51 up at AA143493 310 pleckstrin 2 (mouse) homolog 4.58 up at AA143763 311 EST 18.09 up at AA146619 312 EST up at AA147084 314 proliferation-associated 2G4, 38kD 4.4 up s_at AA147439 315 EST 5.01 up | rc_AA134985_at | AA134985 | 290 | EST | 12.11 | dn | 0.00318 |
| at AA135871 294 EST at AA135894 295 retinoic acid induced 3 at AA136547 302 EST at AA143493 310 pleckstrin 2 (mouse) homolog at AA143763 311 EST at AA147084 314 proliferation-associated 2G4, 38kD s_at AA147439 315 EST s_at AA147439 315 EST 5.01 up | ¦ | AA135407 | 292 | endogenous retroviral protease | 3.64 | dn | 0.01086 |
| at AA135894 295 retinoic acid induced 3 14.27 up at AA136547 302 EST 3.51 up at AA143493 310 pleckstrin 2 (mouse) homolog 4.58 up at AA143763 311 EST 18.09 up at AA146619 312 EST up at AA147084 314 proliferation-associated 2G4, 38kD 4.4 up s_at AA147439 315 EST 5.01 up | | AA135871 | 294 | EST | 3.39 | dn | 0.01544 |
| at AA136547 302 EST 3.51 up at AA143493 310 pleckstrin 2 (mouse) homolog 4.58 up at AA143763 311 EST 18.09 up at AA146619 312 EST up at AA147084 314 proliferation-associated 2G4, 38kD 4.4 up s_at AA147439 315 EST 5.01 up | | AA135894 | 292 | retinoic acid induced 3 | 14.27 | dn | 0.00558 |
| _at AA143493 310 pleckstrin 2 (mouse) homolog 4.58 up _at AA143763 311 EST _at AA146619 312 EST _at AA147084 314 proliferation-associated 2G4, 38kD 4.44 up _s_at AA147439 315 EST 5.01 up | rc_AA136547_at | AA136547 | 302 | EST | 3.51 | dn | 0.00308 |
| _at AA143763 311 EST 18.09 up | rc_AA143493_at | AA143493 | 310 | pleckstrin 2 (mouse) homolog | 4.58 | dn | 0.01037 |
| _at AA146619 312 EST 4.87 up | | AA143763 | 311 | EST | 18.09 | dn | 0.00347 |
| _at AA147084 314 proliferation-associated 2G4, 38kD 4.4 up s_at AA147439 315 EST 5.01 up | rc_AA146619_at | AA146619 | 312 | EST | 4.87 | dn | 0.00863 |
| AA147439 315 EST 5.01 up | rc_AA147084_at | AA147084 | 314 | proliferation-associated 2G4, 38kD | 4.4 | dn | 0.0045 |
| | rc_AA147439_s_at | AA147439 | 315 | EST | 5.01 | dn | 0.00336 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|----------------|---------|
| rc AA148885 at | AA148885 | 320 | minichromosome maintenance deficient (S. cerevisiae) 4 | 6.39 | Ωn | 0.0125 |
| rc_AA148977_at | AA148977 | 322 | EST | 3.64 | dn | 0.02911 |
| rc_AA149889_at | AA149889 | 326 | neighbor of A-kinase anchoring protein 95 | 3.34 | . da | 0.02054 |
| rc_AA150053_at | AA150053 | 327 | EST | 3.71 | g dn | 0.00102 |
| rc_AA151182_at | AA151182 | 332 | EST | 3.51 | . dn | 0.00043 |
| rc_AA151428_s_at | AA151428 | 335 | matrix metalloproteinase 23B | 7.15 | g. | 0.00056 |
| rc_AA151778_at | AA151778 | 338 | claudin 7 | 5.77 | g. | 0.00002 |
| | | | ATP synthase, H+ transporting, mitochondrial F0 complex, | | • | |
| rc_AA156187_at | AA156187 | 339 | subunit b, isoform 1 | 23.24 | dn | 9000000 |
| rc_AA156243_at | AA156243 | 340 | serine protease, umbilical endothelium | 41.44 | . an | 0.00139 |
| rc_AA156450_at | AA156450 | 342 | EST | 3.33 | . dn | 0.00587 |
| rc_AA157818_f_at | AA157818 | 349 | endogenous retroviral protease | 25.99 | <u> </u> | 0.00153 |
| rc_AA158234_at | AA158234 | 351 | EST | 17.04 | g dn | 0.00711 |
| rc_AA158795_at | AA158795 | 352 | EST | 3.07 | <u>a</u> | 0.00057 |
| rc_AA159525_at | AA159525 | 354 | EST | 49.39 | dn | 0.00062 |
| rc_AA161043_at | AA161043 | 356 | tetraspan 1 | 18.8 | d _n | 0.00015 |
| rc_AA161292_s_at | AA161292 | 357 | interferon, alpha-inducible protein 27 | 13.84 | g, | 0.00004 |
| rc_AA164252_f_at | AA164252 | 358 | VGF nerve growth factor inducible | 3.41 | d d | 0.00154 |
| | | | NADH dehydrogenase (ubiquinone) Fe-S protein 6 (13kD) | | | |
| rc_AA169837_at | AA169837 | 364 | (NADH-coenzyme Q reductase) | 3.17 | dn | 0.03038 |
| rc_AA171760_at | AA171760 | 367 | EST | 17.86 | dn | 0 |
| rc_AA171939_at | AA171939 | 368 | EST | 4.67 | d d | 0.00104 |
| rc_AA172076_at | AA172076 | 369 | EST | 3.06 | dn | 0.00326 |
| rc_AA173430_at | AA173430 | 371 | EST | 4.32 | dn | 0.04362 |
| rc_AA179298_at | AA179298 | 378 | stomatin-like protein 2 | 3.72 | dn | 0.00299 |
| rc_AA179787_at | AA179787 | 380 | polyglutamine binding protein 1 | 6.44 | dn | 0.00206 |
| rc_AA179845_at | AA179845 | 381 | EST | 5.77 | dn | 0.01414 |
| rc_AA181600_at | AA181600 | 384 | EST | 5.38 | dn | 0.03316 |
| rc_AA182001_i_at | AA182001 | 386 | EST | 3.56 | dn | 0.00945 |
| rc_AA187938_at | AA187938 | 391 | EST | 3.86 | dn | 0.00512 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| rc_AA188378_i_at | AA188378 | 392 | | 9.56 | dn | 0.00271 |
| rc_AA188378_f_at | AA188378 | 392 | EST | 5.54 | d d | 0.01359 |
| rc_AA189015_at | AA189015 | 394 | EST . | 5.27 | dn | 0.00004 |
| rc_AA191708_at | AA191708 | 400 | EST | 3.87 | d d | 0.01904 |
| rc_AA192755_at | AA192755 | 401 | EST | 3.13 | dn | 0.00442 |
| rc_AA194237_at | AA194237 | 408 | EST | 3.13 | dn | 0.00212 |
| rc_AA194724_at | AA194724 | 409 | endonuclease G | 3.09 | dh | 0.04011 |
| rc_AA196790_at | AA196790 | 421 | EST | 7.64 | dn | 0.00287 |
| rc_AA204927_at | AA204927 | 425 | tropomyosin 1 (alpha) | 7.09 | dn | 0.00074 |
| rc_AA211483_at | AA211483 | 435 | EST | 44.07 | dn | 0.00175 |
| rc_AA211851_f_at | AA211851 | 436 | EST | 3.66 | dn | 0.00789 |
| rc_AA213696_at | AA213696 | 437 | poly(A)-binding protein, cytoplasmic 1 | 24.65 | dn | 0.00001 |
| AA215299_s_at | AA215299 | 439 | U6 snRNA-associated Sm-like protein LSm7 | 15.84 | dn | 0.00001 |
| rc_AA215468_s_at | AA215468 | 44 | ADP-ribosylation factor-related protein 1 | 3.9 | dn | 0.04549 |
| rc_AA218663_at | AA218663 | 444 | acid-inducible phosphoprotein | 3.26 | dn | 0.03537 |
| AA224502_at | AA224502 | 451 | EST | 4.12 | dn | 0.00694 |
| rc_AA226932_at | AA226932 | 453 | DKFZP564F0923 protein | 6.84 | dn | 0.00405 |
| rc_AA227560_at | AA227560 | 458 | EST | 12.81 | d d | 0.01693 |
| rc_AA227926_at | AA227926 | 460 | EST | 6.81 | dn | 0.01701 |
| rc_AA233886_s_at | AA233886 | 475 | D site of albumin promoter (albumin D-box) binding protein | 3.38 | G | 0.0218 |
| rc_AA233959_i_at | AA233959 | 477 | EST | 19.69 | dn | 0.00101 |
| rc_AA234096_at | AA234096 | 479 | EST | 23.72 | dn | 0.00018 |
| rc_AA234362_at | AA234362 | 481 | EST | 3.89 | dn | 0.03524 |
| rc_AA235707_at | AA235707 | 200 | EST | 9.17 | dn | 0.00005 |
| rc_AA236037_at | AA236037 | 206 | EST | 5.74 | dn | 0.0091 |
| rc_AA236533_s_at | AA236533 | 514 | ecotropic viral integration site 1 | 4.01 | dn | 0.02882 |
| rc_AA236714_f_at | AA236714 | 516 | nuclear mitotic apparatus protein 1 | 4.35 | dn | 0.00083 |
| rc_AA237017_at | AA237017 | 521 | KIAA1068 protein | 3.52 | dn | 0.00976 |
| rc_AA243133_at | AA243133 | 525 | serine/threonine kinase 15 | 7.73 | dn | 0.04328 |
| rc_AA243173_at | AA243173 | 526 | EST | 8.75 | dn | 0.00003 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| AA249819_s_at | AA249819 | 535 | EST | 5.09 | dh | 0.00136 |
| rc_AA251230_at | AA251230 | 540 | EST | 4.75 | dn | 0.00054 |
| rc_AA251299_s_at | AA251299 | 541 | KIAA0014 gene product | 16.86 | dn | 0.00046 |
| rc_AA251909_at | AA251909 | 549 | EST | 3.35 | dn | 0.03937 |
| rc_AA252994_at | AA252994 | 222 | apoptosis inhibitor 4 (survivin) | 3.55 | dn | 0.00075 |
| rc_AA253011_f_at | AA253011 | 558 | KIAA0713 protein | 4.73 | dn | 0.01944 |
| AA253330_s_at | AA253330 | 562 | adaptor-related protein complex 1, gamma 1 subunit | 3.87 | dn | 0.00708 |
| rc_AA253473_at | AA253473 | 267 | EST | 15.23 | dn | 0.00171 |
| rc_AA256273_at | AA256273 | 222 | EST | 4.13 | dn | 0.03874 |
| rc_AA256642_at | AA256642 | 582 | EST | 11.17 | d S | 0.00035 |
| rc_AA258482_s_at | AA258482 | 296 | zinc finger protein | 3.17 | dn | 0.04606 |
| rc_AA261907_at | AA261907 | 603 | DKFZP566E144 protein | 4.54 | dn | 0.02289 |
| rc_AA262477_at | AA262477 | 809 | ribonuclease HI, large subunit | 4.87 | dn | 0.00005 |
| rc_AA262887_at | AA262887 | 610 | EST | 5.4 | dn | 0.04719 |
| rc_AA262943_at | AA262943 | 611 | EST | 13.42 | dn | 0.00234 |
| rc_AA262969_f_at | AA262969 | 613 | ferritin, heavy polypeptide 1 | 4.49 | dn | 0.00013 |
| AA263044_s_at | AA263044 | 615 | H2A histone family, member Y | 4.13 | dn | 0.00024 |
| rc_AA278817_at | AA278817 | 618 | EST | 4.22 | dn | 0.00061 |
| rc_AA278838_s_at | AA278838 | 620 | EST | က | dn | 0.02832 |
| rc_AA279177_at | AA279177 | 624 | lymphocyte antigen 75 | 6.05 | dn | 0.01821 |
| rc_AA279840_at | AA279840 | 632 | titin-cap (telethonin) | 5.58 | dn | 0.01253 |
| rc_AA280283_s_at | AA280283 | 637 | EST | 4.99 | dn | 0.02644 |
| rc_AA280734_i_at | AA280734 | 639 | KIAA0618 gene product | 9.64 | dn | 0.00003 |
| rc_AA280928_at | AA280928 | . 642 | EST | 3.27 | dn | 0.04625 |
| rc_AA282149_s_at | AA282149 | 654 | huntingtin interacting protein-1-related | 4.19 | dn | 0.00091 |
| rc_AA282247_at | AA282247 | 657 | EST | 8 | dn | 0.00014 |
| rc_AA283085_s_at | AA283085 | 299 | EST | 4.93 | dn | 0.00382 |
| rc_AA284879_at | AA284879 | 619 | EST | 24.68 | dn | 0.00001 |
| rc_AA284945_at | AA284945 | 089 | EST | 5.98 | dn | 0.00026 |
| rc_AA287022_s_at | AA287022 | 685 | thymidine kinase 1, soluble | 4.51 | dn | 0.02582 |
| rc_AA287347_at | AA287347 | 687 | EST | 5.47 | dn | 0.00034 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | le Direction | Pvalue |
|------------------|----------|--------|---|-------------|--------------|---------|
| rc_AA287393_at | AA287393 | 989 | EST | 5.68 | dn | 0.00062 |
| rc_AA290674_s_at | AA290674 | 692 | eukaryotic translation initiation factor 4E binding protein 1 | 11.59 | ŝ | 0.00036 |
| rc_AA291137_at | AA291137 | 694 | EST | 5.21 | . <u>q</u> . | 0.00685 |
| rc_AA291139_at | AA291139 | 695 | EST | 8.69 | 9 | 0.04573 |
| rc_AA291168_at | AA291168 | 969 | EST | 40.67 | G | 0.00065 |
| AA291456_s_at | AA291456 | 200 | EST | 3.96 | g | 0.03633 |
| rc_AA291659_at | AA291659 | 702 | EST | 3.83 | g | 0.01934 |
| rc_AA292379_at | AA292379 | 208 | EST | 6.04 | g. | 0.00568 |
| rc_AA292659_at | AA292659 | 710 | EST | 3.48 | d, | 0.00037 |
| rc_AA292765_at | AA292765 | 712 | ZW10 interactor | 7.14 | dn | 0.02623 |
| rc_AA292788_s_at | AA292788 | 714 | EST | 7.69 | d | 0.00967 |
| rc_AA292931_at | AA292931 | 715 | EST | 3.97 | d | 0.00067 |
| rc_AA293719_at | AA293719 | 720 | EST | 4.69 | d | 0.02181 |
| AA295819_s_at | AA295819 | 722 | EST | 80 | g | 0.01793 |
| AA298786_at | AA298786 | 727 | EST | 4.65 | dn | 0.02821 |
| AA306121_at | AA306121 | 729 | EST | 4.85 | dn | 0.00381 |
| AA307748_s_at | AA307748 | 730 | EST | 3.7 | dn | 0.00001 |
| AA320369_s_at | AA320369 | 735 | chromosome 19 open reading frame 3 | 4.33 | dn | 0.00554 |
| AA328993_s_at | AA328993 | 738 | EST | 3.66 | dn | 0.00146 |
| rc_AA331393_at | AA331393 | 739 | EST | 16.73 | dn | 0.00848 |
| rc_AA335091_at | AA335091 | 740 | EST | 5.28 | dn | 0.000 |
| rc_AA335191_f_at | AA335191 | 741 | creatine kinase, brain | 47.35 | dn | 0.00419 |
| rc_AA338729_at | AA338729 | 743 | EST | 3.33 | dn | 0.00046 |
| rc_AA338889_f_at | AA338889 | 745 | actin related protein 2/3 complex, subunit 4 (20 kD) | 10.77 | dn | 0.03782 |
| AA364267_at | AA364267 | 762 | EST | 5.01 | 음 | 0.00255 |
| rc_AA370163_at | AA370163 | 992 | EST | 3.34 | d | 0.00643 |
| rc_AA372018_at | AA372018 | 768 | EST | 14.3 | g | 0.00178 |
| AA372630_s_at | AA372630 | 269 | differentially expressed in hematopoietic lineages | 25.49 | on on | 0.01743 |
| AA384184_s_at | AA384184 | 774 | DKFZP586B0519 protein | 3.38 | g. | 0.01209 |
| rc_AA394121_at | AA394121 | 778 | laminin receptor 1 (67kD, ribosomal protein SA) | 23.78 | dn | 0.00099 |
| • | | | | | | |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|----------------|---------|
| rc_AA397906_at | AA397906 | 782 | DKFZP434I216 protein | 3.2 | 1 - | 0.00138 |
| rc_AA397916_f_at | AA397916 | 784 | EST | 3.27 | t a | 0.02895 |
| rc_AA398205_at | AA398205 | 789 | EST | 4.17 | - S | 0.00004 |
| rc_AA398761_s_at | AA398761 | 799 | EST | 3.43 | . <u>s</u> | 0.00726 |
| rc_AA398908_at | AA398908 | 801 | EST | 38.69 | . д | 0.01089 |
| rc_AA399226_at | AA399226 | 803 | tight junction protein 3 (zona occludens 3) | 3.59 | d d | 0.02002 |
| rc_AA400271_at | AA400271 | 814 | EST | 3.51 | g | 0.00742 |
| rc_AA401958_at | AA401958 | 832 | EST | 3.62 | d d | 0.01232 |
| rc_AA402495_at | AA402495 | 838 | EST | 4.91 | d d | 0.00235 |
| AA402937_at | AA402937 | 843 | EST | 3.11 | dn dn | 0.00182 |
| rc_AA402968_at | AA402968 | 844 | EST | 3.1 | o n | 0.00453 |
| rc_AA403159_at | AA403159 | 845 | Ste-20 related kinase | 7.33 | g. | 0.00187 |
| rc_AA404338_at | AA404338 | 849 | EST | 10.9 | . an | 0.00668 |
| rc_AA405310_at | AA405310 | 856 | EST. | 3.23 | . a | 0.00138 |
| rc_AA405460_at | AA405460 | 857 | EST | 4.02 | an | 0.03492 |
| rc_AA405715_at | AA405715 | 862 | hypothetical protein | 4.68 | - S | 0.00898 |
| rc_AA405791_at | AA405791 | 864 | EST | 21.22 | . a | 0 |
| rc_AA406145_f_at | AA406145 | 870 | EST | 6.71 | . <u>a</u> | 0.00047 |
| rc_AA406218_at | AA406218 | 872 | EST | 4.88 | - <u>a</u> | 0.02194 |
| rc_AA406385_at | AA406385 | 876 | DKFZP564B0769 protein | 3.21 | r an | 0.00724 |
| rc_AA406542_at | AA406542 | 878 | EST | 8.27 | . g | 0.00724 |
| rc_AA410469_at | AA410469 | 883 | EST | 6.3 | dn | 0.00103 |
| rc_AA410508_at | AA410508 | 882 | EST | 16.04 | dn | 0.02635 |
| rc_AA410962_s_at | AA410962 | 887 | peroxisome proliferative activated receptor, delta | 3.45 | dn | 0.04574 |
| rc_AA410972_at | AA410972 | 888 | EST | 3.12 | dn | 0.00023 |
| rc_AA411502_at | AA411502 | 883 | EST | 16.42 | dn | 0.00241 |
| rc_AA411685_at | AA411685 | 830 | EST | 3.83 | dn | 0.00417 |
| rc_AA411813_at | AA411813 | 893 | postmeiotic segregation increased 2-like 11 | 9.76 | dn | 0.03499 |
| rc_AA412301_at | AA412301 | 833 | EST | 4.57 | dn | 0.00026 |
| | AA412403 | 900 | EST | 3.09 | dn | 0.00047 |
| rc_AA412405_s_at | AA412405 | 901 | EST | 13.82 | ф | 0.01021 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|--------------|---------|
| rc_AA412720_at | AA412720 | 905 | EST | 3.6 | dn | 0.01396 |
| rc_AA416963_at | AA416963 | 911 | EST | 3.69 | dn | 0.03956 |
| rc_AA416973_at | AA416973 | 913 | EST | 3.61 | dn | 0.0091 |
| rc_AA419217_at | AA419217 | 923 | DKFZP586E1422 protein | 6.77 | dn | 0.00045 |
| rc_AA421562_at | AA421562 | 934 | anterior gradient 2 (Xenepus laevis) homolog | 56.3 | d, | 0.0041 |
| rc_AA421638_at | AA421638 | 932 | EST | 3.05 | d | 0.00487 |
| rc_AA422049_at | AA422049 | 937 | EST | 3.38 | ď | 0.0067 |
| rc_AA422086_at | AA422086 | 938 | EST | 10.71 | dn | 0.03418 |
| rc_AA422150_at | AA422150 | 939 | cytochrome P540 family member predicted from ESTs | 17.14 | d | 0.00108 |
| rc_AA424029_at | AA424029 | 943 | EST | 89.8 | d | 0.00081 |
| rc_AA424487_at | AA424487 | 945 | EST | 38.41 | g | 0.00002 |
| rc_AA424881_at | AA424881 | 949 | EST | 6.3 | 롸 | 0.00556 |
| rc_AA425279_at | AA425279 | 951 | quiescin Q6 | 6.15 | dn | 0.00083 |
| rc_AA425401_at | AA425401 | 954 | serine/threonine kinase 24 (Ste20, yeast homolog) | 3.22 | dn | 0.00625 |
| ω, | AA425852 | 928 | EST | 7.78 | ф | 0.00239 |
| rc_AA425852_i_at | AA425852 | 928 | EST | 4.8 | dn | 0.03874 |
| rc_AA426447_at | AA426447 | 965 | EST | 4.23 | dn | 0.0309 |
| rc_AA426521_at | AA426521 | 296 | Sjogren's syndrome nuclear autoantigen 1 | 3.47 | dn | 0.01161 |
| rc_AA427442_at | AA427442 | 971 | guanine nucleotide regulatory factor | 3.43 | dn | 0.01547 |
| AA427468_s_at | AA427468 | 973 | claudin 4 | 84.43 | dn | 0 |
| rc_AA427636_at | AA427636 | 916 | EST | 19.23 | dn | 0.00145 |
| rc_AA427825_at | AA427825 | 981 | EST . | 3.32 | d | 0.01615 |
| rc_aa427925_s_at | AA427925 | 982 | EST | 3.23 | dn | 0.01806 |
| rc_AA427946_at | AA427946 | 983 | dynein, axonemal, light polypeptide 4 | 3.01 | dn | 0.00001 |
| AA428172_f_at | AA428172 | 986 | Notch (Drosophila) homolog 3 | 9.53 | dn | 0.02562 |
| rc_AA428964_at | AA428964 | 993 | kallikrein 10 | 21.83 | dn | 0.02324 |
| rc_AA429009_at | AA429009 | 994 | serine protease inhibitor, Kunitz type 1 | 30.04 | dn | 0.00001 |
| rc_AA429470_at | AA429470 | 966 | EST | 3.67 | dn | 0.00782 |
| rc_AA429472_at | AA429472 | 266 | DKFZP434P106 protein | 8.27 | dn | 0.00208 |
| rc_AA429636_at | AA429636 | 1001 | hexokinase 2 | 10.43 | dn | 0.00597 |
| AA429825_at | AA429825 | 1003 | DKFZP566B023 protein | 6.63 | 함 | 0.00032 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | e Direction | Pvalue |
|------------------|----------|--------|--|-------------|----------------|---------|
| rc_AA429890_s_at | AA429890 | 1004 | cisplatin resistance associated | 12.51 | 17 | 0.00053 |
| rc_AA430032_at | AA430032 | 1009 | pituitary tumor-transforming 1 | 16.87 | d d | 0.00659 |
| rc_AA430048_at | AA430048 | 1012 | KIAA0160 protein | 6.27 | dn | 0.00631 |
| rc_AA430674_at | AA430674 | 1018 | EST | 15.11 | dn | 0.00293 |
| rc_AA431719_at | AA431719 | 1025 | EST | 4.25 | dn | 0.00015 |
| rc_AA431776_at | AA431776 | 1027 | EST | 3.24 | dn | 0.01814 |
| rc_AA431873_at | AA431873 | 1028 | EST | 4.03 | d _n | 0.00785 |
| rc_AA432162_at | AA432162 | 1029 | DKFZP586B2022 protein | 3.48 | d d | 0.03851 |
| rc_AA433930_at | AA433930 | 1032 | chondroitin 4-sulfotransferase | 7.68 | dn | 0.02445 |
| rc_AA434418_at | AA434418 | 1036 | KIAA1115 protein | 5.12 | dn | 0.00498 |
| rc_AA435526_s_at | AA435526 | 1037 | transferrin receptor (p90, CD71) | 3.69 | dn | 0.00139 |
| rc_AA435665_at | AA435665 | 1040 | EST | 8.66 | dn | 0.00001 |
| rc_AA436027_at | AA436027 | 1050 | EST | 3.71 | g | 0.03676 |
| rc_AA436473_s_at | AA436473 | 1052 | EST | 3.03 | dn | 0.00133 |
| rc_AA436616_at | AA436616 | 1056 | EST | 3.18 | g | 0.04402 |
| rc_AA437368_at | AA437368 | 1063 | EST | 3.75 | dn | 0.01317 |
| rc_AA437387_s_at | AA437387 | 1064 | EST | 3.81 | dn | 0.01478 |
| rc_AA441911_at | AA441911 | 1066 | EST | 6.14 | dn | 0.00003 |
| AA442054_s_at | AA442054 | 1067 | phospholipase C, gamma 1 (formerly subtype 148) | 16.89 | d | 0.00205 |
| rc_AA442763_at | AA442763 | 1072 | cyclin B2 | 5.09 | dn | 0.02168 |
| rc_AA443271_at | AA443271 | 1073 | KIAA0546 protein | 3.6 | dn | 0.01228 |
| rc_AA443316_s_at | AA443316 | 1075 | v-Ha-ras Harvey rat sarcoma viral oncogene homolog | 4.13 | dn | 0.01729 |
| rc_AA443941_at | AA443941 | 1085 | tumor suppressing subtransferable candidate 1 | 3.57 | dn | 0.01685 |
| rc_AA446949_at | AA446949 | 1096 | EST | 3.41 | d | 0.03411 |
| rc_AA446968_at | AA446968 | 1097 | EST | 3.45 | dn | 0.02232 |
| rc_AA447118_s_at | AA447118 | 1099 | EST | 3.03 | g. | 0.01702 |
| rc_AA447687_at | AA447687 | 1104 | EST | 11.42 | đ | 0.00362 |
| rc_AA447732_at | AA447732 | 1105 | EST | 3.2 | dn | 0.00591 |
| rc_AA447991_at | AA447991 | 1112 | EST | 4.99 | dn | 0.00173 |
| rc_aa449073_s_at | AA449073 | 1117 | EST | 6.89 | dn | 0.01445 |
| rc_AA449122_at | AA449122 | 1119 | EST | 3.65 | dn | 0.00369 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | e Direction | Pvalue |
|------------------|----------|--------|--|-------------|----------------|---------|
| rc_AA449456_at | AA449456 | 1126 | EST | 6.29 | dn | 0.00087 |
| rc_AA449458_at | AA449458 | 1127 | EST | 3.22 | 9 | 0.03098 |
| rc_AA449475_at | AA449475 | 1128 | EST | 3.06 | đ | 0.00291 |
| rc_AA449479_at | AA449479 | 1129 | EST | 3.09 | đ | 0.03495 |
| rc_AA450247_at | AA450247 | 1133 | EST | 5.27 | a n | 0.02833 |
| rc_AA451676_at | AA451676 | 1135 | EST | 14.72 | e e | 0.00056 |
| | | | hepatocellular carcinoma associated protein; breast cancer | | | |
| rc_AA451680_at | AA451680 | 1136 | associated gene 1 | 3.55 | dn | 0.00708 |
| rc_AA451877_at | AA451877 | 1138 | EST | 8.63 | dn | 0.00489 |
| rc_AA452259_at | AA452259 | 1143 | EST | 3.49 | ф | 0.00114 |
| rc_AA452536_at | AA452536 | 1145 | v-ral simian leukemia viral oncogene homolog A (ras related) | 5.6 | 음 | 0.00481 |
| AA452724_at | AA452724 | 1149 | programmed cell death 5 | 7.2 | d | 0.00908 |
| rc_AA453477_at | AA453477 | 1153 | X-prolyl aminopeptidase (aminopeptidase P)-like | 4.23 | đ | 0.0001 |
| rc_AA453783_s_at | AA453783 | 1158 | EST | 6.16 | ď | 0.00167 |
| rc_AA454597_s_at | AA454597 | 1166 | EST | 3.63 | dn | 0.0067 |
| rc_AA454710_at | AA454710 | 1168 | EST | 3.42 | đ | 0.00653 |
| AA454908_s_at | AA454908 | 1171 | KIAA0144 gene product | 6.3 | đ | 0.00539 |
| rc_AA455521_s_at | AA455521 | 1178 | E2F transcription factor 5, p130-binding | 4.6 | đ | 0.00773 |
| rc_AA455522_s_at | AA455522 | 1179 | EST | 3.83 | đ | 0.00017 |
| rc_aa458852_f_at | AA458852 | 1203 | KIAA0440 protein | 3.2 | đ | 0.00038 |
| rc_AA458890_at | AA458890 | 1206 | EST | 3.36 | g. | 0.00303 |
| rc_AA459254_at | AA459254 | 1211 | EST | 5.36 | ᅀ | 0.0259 |
| rc_AA459310_r_at | AA459310 | 1214 | EST | 3.45 | ф | 0.00179 |
| rc_AA459388_s_at | AA459388 | 1215 | copine I | 5.23 | ф | 0.00691 |
| rc_AA459703_at | AA459703 | 1222 | v-myc avian myelocytomatosis viral oncogene homolog | 4.75 | 유 | 0.02413 |
| rc_AA459961_at | AA459961 | 1223 | EST | 3.24 | đ | 0.00316 |
| rc_AA460017_i_at | AA460017 | 1225 | EST | 10.76 | 육 | 0.00106 |
| rc_AA460017_f_at | AA460017 | 1225 | EST | 3.61 | 육 | 0.00109 |
| rc_AA461187_at | AA461187 | 1236 | EST | 8.15 | ф | 0.00068 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | e Direction | Pyalife |
|--------------------|------------|--------|---|-------------|---------------------------|---------|
| rc_AA461473_at | AA461473 | 1242 | nebulette | 3.35 | | 0.03855 |
| rc_AA461476_at | AA461476 | 1243 | EST | 4.12 | d G | 0.00871 |
| rc_AA463234_at | AA463234 | 1246 | KIAA0792 gene product | 5.94 | ďn | 0.01182 |
| | | | mannosyi (alpha-1,6-)-glycoprotein beta-1,2-N- | | • | |
| rc_AA463725_at | , AA463725 | 1249 | acetylglucosaminyltransferase | 7.4 | dn | 0.04715 |
| rc_AA463861_at | AA463861 | 1251 | EST | 24.79 | g. | 0.00096 |
| . rc_AA464414_i_at | AA464414 | 1258 | EST | 4.99 | d _n | 0.00529 |
| rc_AA464698_at | AA464698 | 1262 | EST | 3.24 | - <u>G</u> | 0.04854 |
| rc_AA464963_at | AA464963 | 1265 | EST | 5.01 | - <u>9</u> | 0.00107 |
| | | | BUB3 (budding uninhibited by benzimidazoles 3, yeast) | | - | |
| AA471278_at | AA471278 | 1277 | homolog | 5.14 | dn | 0.00873 |
| rc_AA476216_at | AA476216 | 1279 | EST | 4.97 | . an | 0.00359 |
| rc_AA478017_at | AA478017 | 1295 | zyxin | 5.77 | . dn | 0.00484 |
| rc_AA478300_at | AA478300 | 1298 | CD39-like 2 | 6.15 | dn | 0.01625 |
| rc_AA478415_at | AA478415 | 1299 | EST | 4.56 | g. | 0.00095 |
| rc_AA478599_at | AA478599 | 1304 | G protein-coupled receptor 56 | 3.31 | d _D | 0.00182 |
| rc_AA479044_s_at | AA479044 | 1307 | EST | 6.9 | dn | 0.04668 |
| rc_AA479727_s_at | AA479727 | 1315 | EST | 90.9 | <u>a</u> | 0.00389 |
| rc_AA479797_at | AA479797 | 1316 | EST | 7.93 | dn | 900000 |
| rc_AA479945_s_at | AA479945 | 1319 | plakophilin 3 | 3.17 | dn | 0.01767 |
| rc_AA482007_at | AA482007 | 1331 | EST | 3.49 | dn | 0.00167 |
| | | | protein kinase related to S. cerevisiae STE20, effector for | | | |
| rc_AA482127_at | AA482127 | 1333 | Cdc42Hs | 4.88 | <u>a</u> | 0.00017 |
| rc_AA482224_f_at | AA482224 | 1334 | putative type II membrane protein | 4.62 | - S | 0.0105 |
| AA482319_f_at | AA482319 | 1335 | putative type II membrane protein | 5.11 | dn | 0.00177 |
| rc_AA482546_s_at | AA482546 | 1336 | KIAA0124 protein | 4.41 | d _n | 0.00604 |
| rc_AA482613_at | AA482613 | 1338 | DKFZP434B203 protein | 4.14 | g. | 0.00186 |
| rc_AA485405_at | AA485405 | 1343 | EST | 5.35 | d | 0.03475 |
| ធ | AA485697 | 1346 | EST | 14.74 | dn | 0.00102 |
| rc_AA488987_s_at | AA488987 | 1365 | synaptogyrin 2 | 3.24 | dn | 0.01444 |
| rc_AA489707_at | AA489707 | 1371 | EST | 3.47 | dn | 0.03433 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | te Direction | Pvalue |
|------------------|----------|--------|---|-------------|--------------|---------|
| rc_AA489712_at | AA489712 | 1372 | EST | 4.93 | dn | 0.00726 |
| rc_AA490212_at | AA490212 | 1375 | H2A histone family, member Y | 3.71 | d S | 0.01226 |
| rc_AA490494_at | AA490494 | 1377 | EST | 5.16 | 음 | 0.01696 |
| rc_AA491223_at | AA491223 | 1389 | EST | 3.03 | dn | 0.00557 |
| rc_AA496204_at | AA496204 | 1397 | EST | 3.69 | d | 0.01097 |
| rc_AA496245_at | AA496245 | 1398 | EST | 3.96 | d d | 0.0039 |
| | | | v-erb-b2 avian erythroblastic leukemia viral oncogene homolog | | • | |
| rc_AA496981_at | AA496981 | 1404 | | 4.93 | dn | 0.01096 |
| rc_AA497031_at | AA497031 | 1407 | EST | 11.05 | g. | 0.04381 |
| rc_AA504111_at | AA504111 | 1409 | EST | 3.2 | ď | 0.00544 |
| rc_AA504264_at | AA504264 | 1410 | EST | 3.81 | dn | 0.00684 |
| rc_AA504270_at | AA504270 | 1411 | EST | 4.96 | g | 0.01919 |
| AA504413_at | AA504413 | 1413 | EST | 3.35 | d. | 0.00079 |
| rc_AA504806_at | AA504806 | 1416 | EST | 3.54 | dn | 0.00221 |
| rc_AA598405_at | AA598405 | 1424 | membrane interacting protein of RGS16 | 4.69 | dn | 0.0122 |
| rc_AA598506_s_at | AA598506 | 1430 | KIAA0179 protein | 3.17 | d | 0.01694 |
| rc_AA598712_at | AA598712 | 1436 | EST | 3.03 | dn | 0.03656 |
| rc_AA598988_at | AA598988 | 1442 | EST | 4.32 | g | 0.00044 |
| rc_AA599244_at | AA599244 | 1448 | KIAA0530 protein | 3.39 | dn | 0.01246 |
| rc AA599522 f at | AA599522 | 1452 | squamous cell carcinoma antigen recognised by T cells | 6.75 | 9 | 0.04229 |
| rc_AA608579_s_at | AA608579 | 1464 | paired-like homeodomain transcription factor 2 | 4.29 | . <u>a</u> | 0.04435 |
| rc_AA608897_at | AA608897 | 1473 | EST | 9.92 | g | 0.00087 |
| rc_AA608965_at | AA608965 | 1474 | Hermansky-Pudlak syndrome | 3.19 | d | 0.00204 |
| rc_AA609008_at | AA609008 | 1475 | EST | 3.46 | dn | 0.02935 |
| rc_AA609013_s_at | AA609013 | 1477 | dipeptidase 1 (renal) | 10.17 | dn | 0.00109 |
| rc_AA609614_at | AA609614 | 1487 | EST | 6.5 | d n | 0.00406 |
| rc_AA609786_s_at | AA609786 | 1491 | nucleolar protein 1 (120kD) | 4.75 | đ | 0.00261 |
| rc_AA610053_at | AA610053 | 1496 | EST | 7.01 | dn | 0.00003 |
| rc_AA610116_i_at | AA610116 | 1499 | tetraspan NET-6 protein | 33.68 | dn | 0.00171 |
| rc_AA620466_at | AA620466 | 1502 | EST | 5.14 | dn | 0.00004 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| rc_AA620881_at | AA620881 | 1510 | trinucleotide repeat containing 3 | 8.66 | dn | 0.00735 |
| | AA620995 | 1512 | EST | 3.74 | ф | 0.03414 |
| rc_AA621277_at | AA621277 | 1520 | EST | 3.81 | dn | 0.00194 |
| rc_AA621780_at | AA621780 | 1530 | CGI-96 protein | 3.65 | dn | 0.01582 |
| AB000584_at | AB000584 | 1533 | prostate differentiation factor | 4.7 | dn | 0.00071 |
| AB002533_at | AB002533 | 1539 | karyopherin alpha 4 (importin alpha 3) | 6.38 | dn | 0.00003 |
| AB006781_s_at | AB006781 | 1540 | lectin, galactoside-binding, soluble, 4 (galectin 4) | 7.05 | dn | 0.00913 |
| AF001294_at | AF001294 | 1544 | tumor suppressing subtransferable candidate 3 | 7.45 | dn | 0.0000 |
| AF003521_at | AF003521 | 1545 | • | 11.26 | ф | 0.00008 |
| AF004709_at | AF004709 | 1547 | mitogen-activated protein kinase 13 | 3.92 | ф | 0.0009 |
| | | | stress-associated endoplasmic reticulum protein 1; ribosome | | | |
| C00021_s_at | C00021 | 1551 | associated membrane protein 4 | 3.33 | d | 0.00215 |
| C01766 s at | C01766 | 1559 | | 13.67 | d | 0.00003 |
| rc_C13992_f_at | C13992 | 1564 | EST | 6.39 | g | 0.00059 |
| rc_C14051_f_at | C14051 | 1565 | phosphoprotein enriched in astrocytes 15 | 3.68 | ġ | 0.01453 |
| rc C14098 f at | C14098 | 1566 | EST | 3.53 | dn | 0.04401 |
| rc C14348 at | C14348 | 1568 | EST | 4.06 | dn | 0.00111 |
| C14412 s at | C14412 | 1569 | HSPC038 protein | 3.9 | dn | 0.00036 |
| rc_C14756_f_at | C14756 | 1570 | MLN51 protein | 5.36 | dn | 0.00001 |
| rc_C15324_f_at | C15324 | 1574 | _ | 5.22 | dn | 0.00344 |
| rc_C21248_at | C21248 | 1585 | pituitary tumor-transforming 1 | 3.85 | ďn | 0.00456 |
| D00017_at | D00017 | 1587 | annexin A2 | 11.38 | dn | 0 |
| | | | interleukin 2 receptor, gamma (severe combined | | | |
| D11086_at | D11086 | 1595 | | 5.61 | d d | 0.00873 |
| | | | heterogeneous nuclear ribonucleoprotein U (scaffold | | | |
| D13413 rna1 s_at | D13413 | 1604 | attachment factor A) | 4.79 | ф | 0.00092 |
| D13639_at | D13639 | 1607 | cyclin D2 | 7.49 | dn | 0.01641 |
| D14520_at | D14520 | 1613 | basic transcription element binding protein 2 | 4.93 | dn | 0.00004 |
| D14530_at | D14530 | 1614 | ribosomal protein S23 | 3.1 | dn | 0.00331 |
| D14657_at | D14657 | 1615 | KIAA0101 gene product | 3.7 | dn | 0.04079 |
| rc_D19737_at | D19737 | 1623 | golgi autoantigen, golgin subfamily a, 3 | 3.44 | ф | 0.02212 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | • Direction | Pvalue |
|----------------|---------|--------|---|-------------|-------------|----------|
| rc_D20464_at | D20464 | 1625 | | 3.27 | dn | 0.04897 |
| rc_D20906_at | D20906 | 1627 | EST | 5.18 | dn | 0.02189 |
| _ | | | minichromosome maintenance deficient (S. cerevisiae) 2 | | | |
| D21063_at | D21063 | 1628 | (mitotin) | 3.83 | dn | 0.00983 |
| D21261_at | D21261 | 1629 | transgelin 2 | 3.46 | ф | 0.00685 |
| D23660_at | D23660 | 1630 | ribosomal protein L4 | 3.22 | ф | 0.00316 |
| D25216_at | D25216 | 1631 | KIAA0014 gene product | 3.17 | d | 0.02125 |
| D25274_at | D25274 | 1632 | EST | 3.39 | dn | 0.00238 |
| D25328_at | D25328 | 1633 | phosphofructokinase, platelet | 3.17 | dn | 0.04925 |
| rc_D25560_i_at | D25560 | 1634 | EST | 4.72 | dn | 0.00661 |
| D26129_at | D26129 | 1635 | ribonuclease, RNase A family, 1 (pancreatic) | 5.68 | d | 0.03827 |
| | | | neuroblastoma candidate region, suppression of tumorigenicity | | • | |
| D28124_at | D28124 | 1636 | ν- | 6.38 | dn | 0.00015 |
| D31094_at | D31094 | 1639 | G8 protein | 4.41 | d | 0.04845 |
| D31417_at | D31417 | 1645 | secreted protein of unknown function | 4.56 | dn | 0.00014 |
| | | | | | | |
| D38073_at | D38073 | 1651 | minichromosome maintenance deficient (S. cerevisiae) 3 | 3.65 | dn | 0.01869 |
| D38548_at | D38548 | 1655 | KIAA0076 gene product | 4.5 | ф | 0.00036 |
| D38583_at | D38583 | 1656 | S100 calcium-binding protein A11 (calgizzarin) | 18.7 | dn | 0.00003 |
| D42085_at | D42085 | 1658 | KIAA0095 gene product | 3.83 | dn | 0.00036 |
| D43949_at | D43949 | 1659 | KIAA0082 protein | 3.32 | dn | 0.0014 |
| D43950_at | D43950 | 1660 | chaperonin containing TCP1, subunit 5 (epsilon) | 3.61 | dn | 0.00672 |
| D49400_at | D49400 | 1667 | ATPase, vacuolar, 14 kD | 4.01 | dn | 0.00287 |
| D50663_at | D50663 | 1671 | t-complex-associated-testis-expressed 1-like 1 | 4.09 | dn | 0.00208 |
| D50913_at | D50913 | 1672 | KIAA0123 protein | 3.32 | dn | 0.01202 |
| D50914_at | D50914 | 1673 | KIAA0124 protein | 4.74 | dn | 0,00752. |
| rc_D51112_s_at | D51112 | 1675 | collapsin response mediator protein 1 | 10.48 | dn | 0.00076 |
| rc_D51133_f_at | D51133 | 1676 | tubulin, beta, 4 | 4.84 | dn | 0.02875 |
| rc_D51276_f_at | D51276 | 1678 | leukemia-associated phosphoprotein p18 (stathmin) | 4.81 | dn | 0.00514 |
| rc_D51287_f_at | D51287 | 1680 | ribosomal protein S12 | 3.3 | dn | 0.02829 |
| rc_D51393_f_at | D51393 | 1681 | ribosomal protein L4 | 5.64 | ф | 0.00074 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|-------------|---|-------------|------------------|---------|
| rc_D52632_f_at | D52632 | 1683 | ribosomal protein S6 | 3.12 | an | 0.00498 |
| rc_D53139_f_at | D53139 | 1684 | ribosomal protein S28 | 4.38 | - 9 | 0.000 |
| rc_D54296_f_at | D54296 | 1685 | KIAA0255 gene product | 3.8 | d d | 0.01059 |
| D55716_at | D55716 | 1686 | minichromosome maintenance deficient (S. cerevisiae) 7 | 6.6 | 9 | 0.00151 |
| rc_D57489_at | D57489 | 1689 | chaperonin containing TCP1, subunit 7 (eta) | 4.08 | î S | 0.00001 |
| rc_D59322_f_at | D59322 | 1694 | EST | 4.18 | - a n | 0.00000 |
| rc_D59847_at | D59847 | 1701 | EST | 3.09 | . <u>a</u> | 0.02206 |
| rc_D60811_s_at | D60811 | 1704 | EST | 3.22 | . g | 0.0098 |
| | | | platelet-activating factor acetylhydrolase, isoform lb, gamma | | | |
| D63391_at | D63391 | 1710 | subunit (29kD) | 3.91 | dn | 0.00062 |
| D63486_at | D63486 | 1712 | KIAA0152 gene product | 6.33 | dn | 0.00078 |
| D63487_at | D63487 | 1713 | KIAA0153 protein | 5.4 | d | 0.00027 |
| D69074 ot | 70000 | 1711 | | č | | |
| D030/4_at | D03074 | 51 / | riigii-iiiobiiity group (nonnistone chromosomai) protein 1 | 3.64 | d | 0.00228 |
| D63880_at | D63880 | 1715 | KIAA0159 gene product | 4.26 | d | 0.00253 |
| D78361_at | D78361 | 1718 | EST | 3.14 | d n | 0.00023 |
| D78676_at | D78676 | 1719 | EST | 3.2 | dn | 0.00635 |
| D79205_at | D79205 | 1721 | ribosomal protein L39 | 3.07 | o S | 0.00021 |
| rc_D80237_s_at | D80237 | 1729 | actin related protein 2/3 complex, subunit 4 (20 kD) | 3.27 | dn dn | 0.00137 |
| rc_D80662_s_at | D80662 | 1733 | adaptor-related protein complex 1, gamma 2 subunit | 3.43 | g. | 0.00108 |
| rc_D80710_f_at | D80710 | 1734 | integral type I protein | 7.08 | dn | 0.00213 |
| rc_D80917_f_at | D80917 | 1736 | KIAA0670 protein/acinus | 3.58 | D | 0.00007 |
| rc_D80946_f_at | D80946 | 1737 | SFRS protein kinase 1 | 8.53 | g | 0.00455 |
| | | | 5-aminoimidazole-4-carboxamide ribonucleotide | | • | |
| D82348_at | D82348 | 1744 | formyltransferase/IMP cyclohydrolase | 3.93 | dn | 0.00037 |
| D82558_at | D82558 | 1746 | novel centrosomal protein RanBPM | 5.94 | dn | 0.00752 |
| D83735_at | D83735 | 1747 | calponin 2 | 10.42 | dn | 0.00001 |
| | , | | trinucleotide repeat containing 11 (THR-associated protein, 230 | | | |
| D83783_at | D83783 | 1748 | kDa subunit) | 6.55 | dn | 0.00176 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq (D | Known Gene Name | Fold Change | e Direction | Pvalue |
|----------------|---------|--------|---|-------------|--------------|---------|
| | | | trinucleotide repeat containing 11 (THR-associated protein, 230 | | | |
| D83783_at | D83783 | 1748 | kDa subunit) | 3.33 | g | 0.00748 |
| D86956_at | D86956 | 1753 | heat shock 105kD | 4 | g | 0.03013 |
| D86974_at | D86974 | 1756 | KIAA0220 protein | 3.85 | g | 0.0161 |
| D87735_at | D87735 | 1764 | ribosomal protein L14 | 3.92 | g | 0.00015 |
| D87953_at | D87953 | 1765 | N-myc downstream regulated | 6.12 | dn | 0.00033 |
| D88154_at | D88154 | 1766 | villin-like | 4.18 | d | 0.00051 |
| rc_F01444_f_at | F01444 | 1770 | KIAA0440 protein | 6.78 | g | 0.00028 |
| rc_F01568_at | F01568 | 1772 | EST | 3.24 | g. | 0.0018 |
| rc_F02800_at | F02800 | 1780 | EST | 3.45 | dn | 0.03238 |
| rc_F02863_at | F02863 | 1782 | EST | 3.21 | S | 0.01039 |
| rc_F04320_s_at | F04320 | 1786 | replication factor C (activator 1) 4 (37kD) | 3.63 | d | 0.01119 |
| rc_F04444_at | F04444 | 1788 | EST | 4.57 | dn | 0.01132 |
| rc_F04531_s_at | F04531 | 1791 | Kell blood group precursor (McLeod phenotype) | 7.79 | ဌ | 0.03205 |
| rc_F04674_at | F04674 | 1793 | KIAA0746 protein | 8.2 | ġ | 0.00028 |
| rc_F09297_s_at | F09297 | 1800 | EST | 3.94 | dn | 0.0016 |
| ທຸ | F09394 | 1803 | KIAA0715 protein | 22.89 | 음 | 0.01753 |
| rc_F09684_at | F09684 | 1805 | EST | 3.78 | d | 0.00277 |
| | | | procollagen-proline, 2-oxoglutarate 4-dioxygenase (proline 4- | | | |
| rc_F09788_at | F09788 | 1808 | hydroxylase), alpha polypeptide II | 4.14 | dn | 0.00003 |
| rc_F13809_f_at | F13809 | 1828 | tropomyosin 1 (alpha) | 7.62 | dn | 0.00012 |
| rc_H04753_f_at | H04753. | 1839 | EST | 3.38 | dn | 0.02447 |
| rc_H04799_at | H04799 | 1841 | EST | 3.71 | d | 0.04109 |
| rc_H05394_f_at | H05394 | 1845 | KIAA0266 gene product | 4.06 | dn | 0.0015 |
| rc_H05525_s_at | H05525 | 1846 | hypothetical protein | 4.6 | dn | 0.0033 |
| ب | H05625 | 1847 | EST | 5.17 | dn | 0.04551 |
| rc_H08863_at | H08863 | 1859 | hypothetical protein | 3.48 | dn | 0.00205 |
| rc_H09241_s_at | H09241 | 1861 | EST | 4.17 | dn | 0.00727 |
| rc_H09271_f_at | H09271 | 1862 | EST | 5.06 | dn | 0.0016 |
| rc_H09281_at | H09281 | 1863 | EST | 9 | dn | 0.00966 |
| rc_H13532_f_at | H13532 | 1881 | ribosomal protein L18a | 3.97 | dn | 0.00061 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|--------------|---------|
| rc_H17476_at | H17476 | 1889 | EST | 3.44 | dn | 0.00479 |
| rc_H18412_s_at | H18412 | 1890 | isocitrate dehydrogenase 3 (NAD+) gamma | 4.37 | dn | 0.00262 |
| rc_H18442_f_at | H18442 | 1891 | creatine kinase, brain | 17.42 | dn | 0.02391 |
| rc_H20989_s_at | H20989 | 1899 | pyruvate kinase, muscle | 11.37 | dn | 0.0375 |
| rc_H24077_at | H24077 | 1900 | EST | 3.05 | dn | 0.0324 |
| rc_H27188_f_at | H27188 | 1908 | collagen-binding protein 2 (colligen 2) | 4.16 | dn | 0.02073 |
| rc H28333 f at | H28333 | 1912 | melanoma adhesion molecule | 3.11 | dn | 0.00172 |
| rc_H29565_at | H29565 | 1913 | EST | 3.89 | dn | 0.01856 |
| rc_H38240_s_at | H38240 | 1916 | thrombospondin 2 | 4.28 | dn | 0.0066 |
| rc_H38568_s_at | H38568 | 1918 | EST | 4.86 | dn | 0.0003 |
| rc_H42321 f at | H42321 | 1928 | ribosomal protein L18a | 3.23 | dn | 0.01102 |
| H43286_s_at | H43286 | 1929 | gamma-aminobutyric acid (GABA) B receptor, 1 | 5.02 | dn | 0.01972 |
| rc_H43646_at | H43646 | 1930 | H2A histone family, member Y | 4.6 | dn | 0.00147 |
| H46486 s at | H46486 | 1932 | nesca protein | 4.77 | dn | 0.00421 |
| rc_H49637_s_at | H49637 | 1940 | EST | 3.79 | dn | 0.01092 |
| rc_H52673_s_at | H52673 | 1943 | BCL2-antagonist/killer 1 | 3.03 | dn | 0.0393 |
| H53657_s_at | H53657 | 1945 | adenylate cyclase 3 | 3.98 | dn | 0.0045 |
| H55437_at | H55437 | 1948 | kraken-like | 3.53 | dn | 0.02344 |
| rc_H56345_r_at | H56345 | 1950 | EST | 4.15 | dn | 0.00488 |
| ! . | | | solute carrier family 2 (facilitated glucose transporter), member | | | |
| rc_H58873_s_at | H58873 | 1961 | _ | 57.98 | dn | 0.00063 |
| rc_H59617_at | H59617 | 1964 | EST | 3.3 | dn | 0.04588 |
| rc H75933 f at | H75933 | 1998 | laminin receptor 1 (67kD, ribosomal protein SA) | 5.81 | dn | 0.00024 |
| rc_H78211_at | H78211 | 2001 | EST | 6.73 | dn | 0.02488 |
| rc_H78323_at | H78323 | 2002 | transcription factor Dp-1 | 3.69 | d | 0.00326 |
| | | | high-mobility group (nonhistone chromosomal) protein isoforms | | | |
| rc_H81413_f_at | H81413 | 2007 | I and Y | 5.82 | ф | 0.00769 |
| rc_H88674_s_at | H88674 | 2021 | collagen, type I, alpha 2 | 5.06 | dn | 0.00866 |
| H89551_s_at | H89551 | 2024 | EST | 9.94 | dn | 0.00137 |
| rc_H93021_at | H93021 | 2033 | peptidylprolyl isomerase A (cyclophilin A) | 3.31 | dn | 0.0183 |
| rc_H93492_at | H93492 | 2037 | EST | 3.94 | ф | 0.01136 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change Direction Pvalue | Direction | Pvalue |
|----------------|---------|--------|---|------------------------------|----------------|---------|
| rc_H93652_f_at | H93652 | | 2039 ribosomal protein S5 | 3.31 | dn | 0.00788 |
| rc_H94471_at | H94471 | | 2042 occludin | 43.09 | g | 0 |
| rc_H95233_s_at | H95233 | 2048 | serine protease inhibitor, Kunitz type, 2 | 47.06 | . д | 0 |
| rc_H96975_at | H96975 | 2057 | EST | 3.22 | d d | 0.0141 |
| rc_H97013_at | H97013 | 2059 | ephrin-A4 | 9.14 | g. | 0.00346 |
| rc_H97809_at | H97809 | 2063 | EST | 4.05 | g. | 0.00111 |
| rc_H98924_at | H98924 | 2072 | chromatin assembly factor 1, subunit A (p150) | 3.53 | . | 0.02106 |
| rc_H99473_s_at | H99473 | 2077 | 2077 regulator of nonsense transcripts 1 | 5.37 | g. | 0.00177 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|---------|--------|---|-------------|-----------|---------|
| | | | | | | |
| J03459 at | J03459 | 2093 | leukotriene A4 hydrolase | 3.03 | dn | 0.04041 |
| J03464 s at | J03464 | 2094 | collagen, type I, alpha 2 | 9.81 | dn | 0.00028 |
| l I | | | solute carrier family 25 (mitochondrial carrier; adenine | | | |
| J03592_at | J03592 | 2096 | nucleotide translocator), member 6 | 10.05 | dn | 0 |
| J03827_at | J03827 | 2100 | nuclease sensitive element binding protein 1 | 4.45 | dn | 0.00015 |
| | | | membrane component, chromosome 1, surface marker 1 | | | |
| J04152 rna1 s at | J04152 | 2107 | (40kD glycoprotein, identified by monoclonal antibody GA733) | 5.26 | dn | 0.02466 |
| J04164 at | J04164 | 2108 | interferon induced transmembrane protein 1 (9-27) | 12.37 | d | 0.00001 |
| AFFX-BioDn-3 at | J04423 | 2109 | EST | 54.11 | dn | 0.02774 |
| AFFX-BioDn-3 at | J04423 | 2109 | EST | 48.05 | ф | 0.02203 |
| AFFX-BioDn-3 at | J04423 | 2109 | EST | 21.46 | ф | 0.04283 |
| AFFX-BioB-3_at | J04423 | 2109 | EST | 5.13 | dn | 0.02791 |
| AFFX-BioB-5_at | J04423 | 2109 | EST | 4.47 | dn | 0.02754 |
| AFFX-BioDn-3 st | J04423 | 2109 | EST | 4.35 | dn | 0.01245 |
| J04469 at | J04469 | 2111 | creatine kinase, mitochondrial 1 (ubiquitous) | 7.9 | dn | 0.00705 |
| J04823 rna1 at | J04823 | 2115 | cytochrome c oxidase subunit VIII | 3.35 | dn | 0.00075 |
| J05257 at | J05257 | 2118 | dipeptidase 1 (renal) | 12.02 | dn | 0.02099 |
| J05582 s at | J05582 | 2121 | mucin 1, transmembrane | 5.39 | dn | 0.00056 |
| ! ! | | | solute carrier family 2 (facilitated glucose transporter), member | | | |
| K03195 at | K03195 | 2128 | | 10.73 | dn | 0.00139 |
| K03460 at | K03460 | 2129 | tubulin, alpha 1 (testis specific) | 7.5 | dn | 0.00002 |
| L03411 s at | L03411 | 2134 | RD RNA-binding protein | 3.97 | dn | 0.00422 |
| L04483 s at | L04483 | 2136 | ribosomal protein S21 | 4.2 | dn | 0.00057 |
| 1 | | | | | | |
| L04490_at | L04490 | 2137 | (39KD) | 5.22 | dn | 0.02192 |
| | | | | | | |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| L06499 L06505 at L08044 L11566 L11669 L11566 L11669 L112350 at L12711 L20941 L20941 L20941 L20941 L20941 L20941 L20941 L23808 L25081 L33075 at L33842 L34587 L34587 | ribosomal protein L37a ribosomal protein L12 trefoil factor 3 (intestinal) trefoil factor 3 (intestinal) | 4 34 | Ē | - |
|---|--|-------|----------|---------|
| L06505 L08044 L08044 L11566 L11669 L12350 L12711 L19527 L19605 at L19686 L20591 L20941 L20591 L2081 L25081 L33075 at L33830 L34587 L34587 | | | 3 | 0.01103 |
| L08044 L08044 L08044 L11566 L11669 L12350 L12350 L1257 L19605 at L19605 L20591 L20591 L2081 L2081 L25081 L33075 at L33930 L34587 L34587 | · · | 10 | dn | 0.00163 |
| L08044 L09604 L11566 L11669 L12350 L12350 L12711 at L19605 L20591 L20591 L20941 L2081 L25081 L33075 at L33930 L34587 L34587 | ٠. | | dn | 0.01674 |
| L09604 L11566 L11669 L12350 L12711 at L19627 L19605 at L20941 L20941 L20941 L20941 L21954 L21954 L21954 L23075 at L33842 L33930 L34587 | | | dn | 0.02124 |
| at L17566 L11669 L12350 L12711 L19627 L19605 at L20941 L20941 L20941 L20941 L21954 L21954 L23081 L33075 at L33842 L33930 L34587 | 2151 proteolipid protein 2 (colonic epithelium-enriched) | 89.8 | dn | 0 |
| at L1711 L12350 L12350 L12711 L19605 at L19686 L20591 L20941 L20941 L21954 L21954 L21954 L21954 L23081 L33075 at L33842 L33930 L33930 | 2156 ribosomal protein L18 | 4.29 | dn | 0.00014 |
| at L12350 L12711 L19527 L19605 at L20591 L20591 L20591 L21954 L21954 L21954 L21954 L23081 L33075 at L33842 L33930 L34587 | 2157 tetracycline transporter-like protein | 6.75 | . dr | 0.00101 |
| at L12711 L19527 L19605 at L19686 L20591 L20941 L21954 L23808 L25081 L33075 at L33842 L3342 L34587 L34587 | 2160 thrombospondin 2 | 3.78 | dn | 0.00061 |
| at L17131 L19605 L19605 at L20591 L20941 L20841 L21954 L23808 L25081 L33075 at L33842 L33930 L34587 | 2161 transketolase (Wernicke-Korsakoff syndrome) | 3.08 | dn | 0.03362 |
| at L17131 L19527 L19605 at L20591 L20591 L20841 L20841 L21954 L23808 L25081 L33075 at L33842 L33930 L34587 | high-mobility group (nonhistone chromosomal) protein isoforms | | | |
| L19527 L19605 L19686 L20591 L20941 L21954 L23808 L25081 L33075 at L33842 L33930 L34587 L34587 | 2168 land Y | 20.57 | dn | 0.00058 |
| at L19605 L20591 L20591 L20941 L21954 L25081 L25081 L33075 at L33842 L33930 L34587 L34587 | 2169 ribosomal protein L27 | | dn dn | 0.00025 |
| a1_at L19686 L20591 L20941 L21954 L23808 L25081 L33075 a1_at L33842 at L33930 L34587 L36720 | 2170 annexin A11 | 6.38 | dn | 0.00017 |
| a1_at L19686 L20591 L20941 L21954 L23808 L25081 L33075 a1_at L33842 at L33930 L34587 L36720 | macrophage migration inhibitory factor (glycosylation-inhibiting | | | |
| L20591 L20941 L21954 L23808 L25081 L33075 a1_at L33842 at L33930 L34587 L34587 | 2171 factor) | 5.26 | dn | 0.00562 |
| L20941 L21954 L23808 L25081 L33075 a1_at L33842 at L33930 L34587 L34587 | 2173 annexin A3 | | dn | 0.00065 |
| L21954 L23808 L25081 L33075 a1_at L33842 at L33930 L34587 L36720 | 2174 ferritin, heavy polypeptide 1 | | dn | 0.01172 |
| L23808 L25081 L33075 a1_at L33842 at L33930 L34587 L36720 | 2177 benzodiazapine receptor (peripheral) | | dn | 0.00001 |
| L25081 L33075 a1_at L33842 at L33930 L34587 L36720 | 2179 matrix metalloproteinase 12 (macrophage elastase) | | dn | 0.02195 |
| L33075 a1_at L33842 at L33930 L34587 L36720 | 2180 ras homolog gene family, member C | 3.67 | dn | 0.00005 |
| a1_at L33842 at L33930 L34587 L36720 | 2195 IQ motif containing GTPase activating protein 1 | 3.83 | dn | 0.00015 |
| at L33930 L34587 L36720 | 2197 IMP (inosine monophosphate) dehydrogenase 2 | 11.03 | dn | 0.00001 |
| L34587 L36720 | 2198 CD24 antigen (small cell lung carcinoma cluster 4 antigen) transcription elongation factor B (SIII), polypeptide 1 (15kD, | 9.16 | 유 | 0.01252 |
| L36720 | 2200 elongin C) | 4.32 | dn | 0.00287 |
| | 2205 bystin-like | - | 슠 | 0.00094 |
| L38696_at L38696 22 | 2208 RNA-binding protein (autoantigenic) thyroid receptor interacting | 3.7 | dn | 0.00093 |
| L40379_at L40379 22 | 2210 protein) | 3.87 | dn | 0.00207 |
| L40904_at L40904 22 | 2212 peroxisome proliferative activated receptor, gamma | 3.43 | g, | 0.03511 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|---------------------------|------------|--------|---|-------------|-----------|---------|
| L41351_at | L41351 | 2214 | protease, serine, 8 (prostasin) | 6.34 | dn | 0.01132 |
| L44538_at | L44538 | 2217 | EST | 4.34 | dn | 0.04319 |
| L76191_at | L76191 | 2222 | interleukin-1 receptor-associated kinase 1 | 5.66 | dn | 0.00089 |
| L76200_at | L76200 | 2223 | guanylate kinase 1 | 3.24 | dn | 0.0097 |
| AFFX-HUMRGE/M1009£M10098 | 39£ M10098 | 2231 | EST | 11.55 | dn | 66000.0 |
| AFFX-HUMRGE/M1009£ M10098 | 39£ M10098 | 2231 | EST | 10.2 | d | 0.00027 |
| AFFX-HUMRGE/M10098 M10098 | 39£ M10098 | 2231 | EST | 5.62 | dn | 0.00251 |
| AFFX-HUMRGE/M10098 M10098 | 39£ M10098 | 2231 | EST | 5.61 | dn | 0.01238 |
| AFFX-HUMRGE/M1009E M10098 | 39£ M10098 | 2231 | EST | 5.35 | dn | 0.00186 |
| AFFX-HUMRGE/M1009£ M10098 | 39£ M10098 | 2231 | EST | 4.79 | dn | 0.00328 |
| AFFX-HUMRGE/M1009£ M10098 | 09£ M10098 | 2231 | EST | 4.66 | dn | 0.00405 |
| AFFX-HUMRGE/M1009£M10098 | 09£ M10098 | 2231 | EST | 3.73 | dn | 0.01794 |
| M12125_at | M12125 | 2241 | tropomyosin 2 (beta) | 10.83 | dn | 0.00191 |
| M13934_cds2_at | M13934 | 2255 | ribosomal protein S14 | 3.99 | dn | 0 |
| M14199_s_at | M14199 | 2258 | laminin receptor 1 (67kD, ribosomal protein SA) | 7.01 | dn | 0 |
| M14483_rna1_s_at | M14483 | 2261 | prothymosin, alpha (gene sequence 28) | 4.46 | dn | 0.00686 |
| M14949_at | M14949 | 2264 | related RAS viral (r-ras) oncogene homolog | 3.11 | dn | 0.00013 |
| M15205_at | M15205 | 2265 | thymidine kinase 1, soluble | 3.75 | dn | 0.00159 |
| M16364_s_at | M16364 | 2269 | creatine kinase, brain | 12.69 | dn | 0.03633 |
| M17733_at | M17733 | 2280 | thymosin, beta 4, X chromosome | 4.15 | dn | 60000.0 |
| M17885_at | M17885 | 2281 | ribosomal protein, large, P0 | 3.92 | dn | 0.00003 |
| M17886_at | M17886 | 2282 | ribosomal protein, large, P1 | 4.34 | d | 0.00004 |
| M18000_at | M18000 | 2283 | ribosomal protein S17 | 3.79 | 롸 | 0.00004 |
| | | | carcinoembryonic antigen-related cell adhesion molecule 6 (non- | | | |
| M18728_at | M18728 | 2285 | specific cross reacting antigen) | 44.82 | dn | 0.00291 |
| M20471_at | M20471 | 2289 | clathrin, light polypeptide (Lca) | 5.32 | dn | 0.00344 |
| M22960_at | M22960 | 2296 | protective protein for beta-galactosidase (galactosialidosis) | 4.49 | dn | 0.00898 |
| M23613_at | M23613 | 2301 | nucleophosmin (nucleolar phosphoprotein B23, numatrin) | 3.67 | dn | 0.00977 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| M24194_at M24485 M24485_s_at M24485 M26708_s_at M27626 M27826_at M27830 AFFX-M27830_5_at M27830 AFFX-M27830_5_at M27830 AFFX-M27830_Mat M27830 AFFX-M27830_Mat M27830 AFFX-M27830_Mat M27830 AFFX-M27830_Mat M27830 AFFX-M27830_Mat M27830 AFFX-M27830_Mat M27830 | 4 2302 5 2304 9314 | guanine nucleotide binding protein (G protein), beta polypeptide | | | |
|--|--------------------------|--|-------|------------|---------|
| व व व व व व व | 000 | | | | |
| , के | .,, | 2-like 1 | 4.55 | dn | 0.00017 |
| # # # # # # # # # # # # # | • | glutathione S-transferase pi | 10.2 | ф | 0.00003 |
| ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,, | • | prothymosin, alpha (gene sequence 28) | 3.35 | ф | 0.00064 |
| | • | endogenous retroviral protease | 26.36 | dn | 0.00342 |
| ्रव, ब्राह्म , ब्राह्म , ब्राह्म , जि. जि. जि. जि. जि. जि. जि. | | EST | 15.53 | dn | 0.00022 |
| | 0 2314 | EST | 14.86 | d | 0.00043 |
| , क , क , क , क , , क , क , क , क , | ., | EST | 10.64 | dn | 0.00213 |
| ,;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;; | ., | EST | 8.6 | dn | 0.00041 |
| , #, #, #, | • | EST | 8.78 | dn | 0.00003 |
| | | EST | 8.2 | dn | 0.00294 |
| , #, | | EST | 6.25 | g. | 0.00046 |
| 1 | ., | EST | 5.78 | dn | 0.0017 |
| M29277_at M29277 | | melanoma adhesion molecule | 3.91 | <u>G</u> . | 0.00112 |
| M29540_at M29540 | .0 2317 | carcinoembryonic antigen-related cell adhesion molecule 5 | 36.57 | dn | 0.0116 |
| M30496 at M30496 | 6 . 2324 | ubiquitin carboxyl-terminal esterase L3 (ubiquitin thiolesterase) | 3.8 | d | 0.00478 |
| a1 at | | leukemia-associated phosphoprotein p18 (stathmin) | 7.48 | dn | 0.00021 |
| s at | | ribosomal protein S24 | 3.37 | dn | 0.00077 |
| € 1 | 0 2328 | ribosomal protein S24 | 3.13 | dn | 0.00014 |
| | 5 2334 | ribosomal protein S15 | 4.01 | dn | 0.00055 |
| M32886_at M32886 | | | 8.35 | d | 0.00215 |
| AFFX-HUMGAPDH/M33 M33197 | | glyceraldehyde-3-phosphate dehydrogenase | 3.31 | dn | 0.00009 |
| M34182 at M34182 | 32 2340 | protein kinase, cAMP-dependent, catalytic, gamma | 4.51 | dn | 0.00043 |
| M35252 at M35252 | 52 2343 | transmembrane 4 superfamily member 3 | 39.12 | ф | 0 |
| M36072_at M36072 | 72 2347 | ribosomal protein L7a S100 calcium-binding protein A10 (annexin II ligand, calpactin I, | 3.1 | dn | 0.00006 |
| M38591_at M38591 | | light polypeptide (p11)) | 14.87 | dn | 0 |
| M38690_at M38690 | 30 2351 | CD9 antigen (p24) | 9.08 | g | 0.00059 |
| M55998_s_at M55998 | 98 2356 | collagen, typę I, alpha 1 | 6.99 | dn | 0.00103 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|--------------|---------|
| M57710 at | M57710 | 2357 | lectin, galactoside-binding, soluble, 3 (galectin 3) | 27.12 | dn | 0.00001 |
| M60784 s at | M60784 | 2366 | small nuclear ribonucleoprotein polypeptide A | 5.74 | dn | 0.00126 |
| M60854 at | M60854 | 2367 | ribosomal protein S16 | 3.4 | dn | 0.00001 |
| M62895 s at | M62895 | 2375 | annexin A2,annexin A2 pseudogene 2 | 6.11 | dn | 0.00013 |
| M64716 at | M64716 | 2382 | ribosomal protein S25 | 3.16 | dn | 0.00039 |
| M68864_at | M68864 | 2389 | ORF | 3.1 | dn | 0.00603 |
| M77232 rna1 at | M77232 | 2399 | ribosomal protein S6 | 3.82 | dn | 0.00045 |
| M77349 at | M77349 | 2400 | transforming growth factor, beta-induced, 68kD | 4.81 | dn | 0.00546 |
| M77836 at | M77836 | 2401 | pyrroline-5-carboxylate reductase 1 | 3.43 | dn | 0.00759 |
| M79463 s at | M79463 | 2402 | promyelocytic leukemia | 4.88 | dn | 0.01821 |
| M81757_at | M81757 | 2406 | ribosomal protein S19 | 5.46 | dn | 0 |
| I | | | tyrosine 3-monooxygenase/tryptophan 5-monooxygenase | | | |
| M86400 at | M86400 | 2410 | activation protein, zeta polypeptide | 5.62 | dn | 0.00016 |
| M86667_at | M86667 | 2411 | nucleosome assembly protein 1-like 1 | 3.03 | dn | 0.04853 |
| l | | | stress-induced-phosphoprotein 1 (Hsp70/Hsp90-organizing | | | |
| M86752_at | M86752 | 2412 | protein) | 7.32 | dn | 0.00001 |
| M87339_at | M87339 | 2415 | replication factor C (activator 1) 4 (37kD) | 4.07 | dn | 0.00316 |
| M91083_at | M91083 | 2419 | chromosome 11 open reading frame 13 | 3.3 | d | 0.00163 |
| 1 | | | membrane component, chromosomal 4, surface marker (35kD | | | |
| M93036_at | M93036 | 2422 | glycoprotein) | 16.45 | d | 0.00308 |
| M94250_at | M94250 | 2426 | midkine (neurite growth-promoting factor 2) | 10.39 | dn | 0.01818 |
| M94345_at | M94345 | 2427 | capping protein (actin filament), gelsolin-like | 22.38 | dn | 0.00003 |
| M96739_at | M96739 | 2434 | nescient helix loop helix 1 | 3.72 | d d | 0.00015 |
| rc_N20198_s_at | N20198 | 2440 | ubiquitin-conjugating enzyme E2 variant 1 | 5.17 | dn | 0.00508 |
| rc N21359 at | N21359 | 2442 | EST | 4.43 | dn | 0.00078 |
| rc_N22015_at | N22015 | 2448 | EST | 46.61 | dn | 0.00025 |
| rc_N22107_at | N22107 | 2449 | EST | 6.88 | dn | 0.04259 |
| rc N24899 at | N24899 | 2461 | EST | 3.06 | dn | 0.00353 |
| rc_N26186_at | N26186 | 2468 | EST | 6.15 | dn | 0.00135 |
| rc N27186 at | N27186 | 2470 | EST | 3.79 | dn | 0.00112 |
| rc_N27334_at | N27334 | 2471 | EST | 3.65 | dn | 0.03437 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affv ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|--------------|---------|
| rc N29888 at | N29888 | 2483 | EST | 3.7 | dn: | 0.00226 |
| rc_N30436_f_at | N30436 | 2484 | EST | 3.45 | dn | 0.02356 |
| rc_N31570_at | N31570 | 2486 | TNF receptor-associated factor 5 | 10.39 | dn | 0.00018 |
| rc_N31597_s at | N31597 | 2487 | DKFZP564G2022 protein | 4.44 | . dn | 0.00085 |
| N36432 at | N36432 | 2507 | erythrocyte membrane protein band 4.1-like 2 | 3.03 | dn | 0.03086 |
| rc N39099 at | M39099 | 2508 | EST | 4.42 | dn | 0.00643 |
| rc_N39237_at | N39237 | 2511 | EST | 9.29 | dn | 0.00001 |
| rc_N39254_s at | N39254 | 2512 | EST | 4.42 | dn | 0.00478 |
| rc_N46423_at | N46423 | 2521 | EST | 9.64 | dn | 0.00027 |
| 1 | | | eukaryotic translation initiation factor 3, subunit 3 (gamma, | , | | |
| rc N47956 at | N47956 | 2524 | 40KD) | 6.34 | dn | 0.00251 |
| rc_N49284_s at | N49284 | 2537 | v-myb avian myeloblastosis viral oncogene homolog | 11.82 | ф | 0.01981 |
| rc_N49738_at | N49738 | 2539 | EST | 3.81 | dn | 0.02479 |
| rc_N50048_at | N50048 | 2542 | EST | 4.08 | dn | 0.00085 |
| rc N51053 s at | N51053 | 2543 | eukaryotic translation initiation factor 5 | 3.42 | dn | 0.01326 |
| rc N51342 at | N51342 | 2545 | EST | 3.56 | dn | 0.0001 |
| rc_N52168_at | N52168 | 2551 | EST | 5.65 | dn | 0.00003 |
| rc_N54841_at | N54841 | 2572 | EST | 42.96 | dn | 0.00002 |
| rc_N56935_s at | N56935 | 2575 | EST | 3.57 | dn | 0.00282 |
| rc_N58463_at | N58463 | 2580 | PCTAIRE protein kinase 1 | 3.18 | dn | 0.00649 |
| rc_N62126_at | N62126 | 2589 | EST | 8.51 | dn | 0.00016 |
| rc_N62675_s at | N62675 | 2594 | DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 16 | 3.61 | dn | 0.04034 |
| rc_N62819_at | N62819 | 2595 | EST | 3.25 | dn | 0.01137 |
| rc_N64616_at | N64616 | 2611 | EST | 3.11 | ф | 0.0074 |
| rc N66139 s at | N66139 | 2615 | neurochondrin | 4.03 | dn | 0.00118 |
| | | | homolog of mouse quaking QKI (KH domain RNA binding | | | |
| rc N66624 at | N66624 | 2618 | protein) | 6.25 | dn | 0 |
| rc_N66951_at | N66951 | 2621 | EST | 5.54 | d | 0.02442 |
| rc_N67205_at | N67205 | 2625 | EST | 3.1 | ф | 0.00626 |
| | | 0 | | 9 75 | <u> </u> | 0.000 |
| rc_N68038_f_at | N68038 | 2632 | phorbolin (similar to apolipoprotein 6 mkina editing protein) | 3.73 | } | - |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| rc N68385 f at | N68385 | 2635 | ribosomal protein L13a | 3.57 | dn | 0.00417 |
| rc_N68921_at | N68921 | 2639 | EST | 4.4 | dn | 0.00989 |
| rc_N69252_f at | N69252 | 2647 | ferritin, light polypeptide | 5.33 | dn | 0.01554 |
| rc_N69263_at | N69263 | 2648 | EST | 7.47 | dn | 0.00004 |
| rc_N70577_at | N70577 | 2659 | EST | 3.07 | dn | 0.01975 |
| rc_N70678_s at | N70678 | 2660 | TAR (HIV) RNA-binding protein 1 | 4.2 | dn | 0.00119 |
| rc N70903 at | N70903 | 2662 | EST | 4.41 | dn | 0.0078 |
| rc_N71072_at | N71072 | 2664 | EST | 2.57 | dn | 0.03881 |
| rc_N71781_at | N71781 | 2666 | | 7.01 | dn | 0.02952 |
| l ŝ | | | solute carrier family 11 (proton-coupled divalent metal ion | | | |
| rc N72116 s at | N72116 | 2668 | transporters), member 2 | 9.01 | dn | 0.00051 |
| rc_N73762_f at | N73762 | 2678 | | 6.65 | dn | 0.0023 |
| rc N73808 f at | N73808 | 2679 | EST | 8.46 | dn | 0.01886 |
| rc N73846 at | N73846 | 2680 | EST | 3.27 | dn | 0.00012 |
| rc N77947 s at | N77947 | 2698 | EST | 2 | dn | 0.00117 |
| rc N80703 at | N80703 | 2704 | EST | 90.9 | dn | 0.00003 |
| rc_N89670_at | N89670 | 2709 | EST | 4.26 | dn | 0.00002 |
| rc_N89937_at | N89937 | 2711 | LIM domain only 7 | 3.6 | dn | 0.00375 |
| rc_N90238 i at | N90238 | 2712 | EST | 3.06 | dn | 0.00354 |
| rc N91023 at | N91023 | 2716 | EST | 3.87 | dn | 0.00008 |
| 1 | | | amyloid beta (A4) precursor protein-binding, family A, member | • | | |
| rc_N92775_at | N92775 | 2723 | 3 (X11-like 2) | 3.86 | dn | 0.00577 |
| rc_N92915_at | N92915 | 2724 | brefeldin A-inhibited guanine nucleotide-exchange protein 1 | 3.1 | <u>d</u> | 0.00807 |
| rc N92934 s at | N92934 | 2725 | cysteine-rich protein 1 (intestinal) | 35.52 | dn | 0.002 |
| rc_N93105_f_at | N93105 | 2728 | EST | 3.77 | dn | 0.02195 |
| rc_N93798_at | N93798 | 2738 | protein tyrosine phosphatase type IVA, member 3 | 4.65 | ф | 0.00118 |
| rc_N98464_s_at | N98464 | 2744 | EST | 15.95 | ф | 0.00004 |
| rc_N98758_f_at | N98758 | 2745 | | 3.87 | dn | 0.0074 |
| N99505_at | N99505 | 2746 | | 3.6 | dn | 0.04499 |
| rc_R02036_at | R02036 | 2754 | EST | 8.01 | dn | 0.01012 |
| | | | | | | |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affv ID | Genbank | Sea ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|----------|--------|--|-------------|-----------|---------|
| rc R06251 f at | R06251 | 2764 | tumor protein D52-like 2 | 5.57 | _ | 0.00037 |
| rc_R06254_f at | R06254 | 2765 | tumor protein D52-like 2 | 4.64 | dn | 0.00039 |
| rc R06866 s at | R06866 | 2774 | EST | 5.18 | ď | 0.00187 |
| rc_R06986_f_at | R06986 | 2776 | peptidylprolyl isomerase B (cyclophilin B) | 3.01 | 음 | 0.04418 |
| rc R22565 at | R22565 | 2800 | EST | 4.8 | dn | 0.0424 |
| rc_R26706_s at | R26706 | 2803 | EST | 3.21 | dn | 0.03858 |
| rc_R26744_at | R26744 | 2804 | midline 1 (Opitz/BBB syndrome) | 4.32 | dn | 0.00532 |
| rc_R27432_at | R27432 | 2808 | EST | 3.62 | dn | 0.00014 |
| i I | | | UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, | | • | |
| rc R28636 at | R28636 | 2809 | polypeptide 3 | 3.78 | dn | 0.00765 |
| rc_R31107_at | R31107 | 2812 | EST | 4.12 | dn | 0.00003 |
| rc_R33498_s at | R33498 | 2820 | EST | 41.34 | dn | 0.00001 |
| rc_R36947 s at | R36947 | 2825 | calcium channel, voltage-dependent, beta 3 subunit | 4.11 | d | 0.00006 |
| rc_r38076 s_at | R38076 | 2828 | EST | 4.08 | dn | 0.00374 |
| rc_R38239_at | R38239 | 2830 | EST | 7.14 | dn | 0.00249 |
| rc_R38280_at | R38280 | 2831 | BCS1 (yeast homolog)-like | 3.68 | dn | 0.0009 |
| rc_R38511 s at | R38511 | 2832 | protein similar to E.coli yhdg and R. capsulatus nifR3 | 5.19 | dn | 0.00015 |
| rc R39191 s at | R39191 | 2834 | KIAA1020 protein | 4.69 | dn | 0.00456 |
| rc_R40254 at | R40254 | 2840 | EST | 5.82 | d | 0.00304 |
| rc R43952 at | R43952 | 2853 | homeo box B5 | 4.11 | dn | 0.04316 |
| rc R44479 at | R44479 | 2855 | KIAA0552 gene product | 4.14 | dn | 0.0181 |
| rc_R44538_at | · R44538 | 2856 | EST | 5.73 | dn | 0.01015 |
| rc_r45698_at | R45698 | 2866 | EST | 3.01 | dn | 0.04766 |
| rc R45994 f at | R45994 | 2867 | EST | 7.81 | dn | 0.0018 |
| rc R48447 at | R48447 | 2871 | EST | 7.75 | dn | 0.00049 |
| rc_R48589_at | R48589 | 2874 | EST | 4.95 | dn | 0.01346 |
| rc_R49084's at | R49084 | 2879 | KIAA0770 protein | 3.57 | dn | 0.00447 |
| rc R49216 at | R49216 | 2880 | EST | 3.64 | d | 0.0004 |
| rc R49395 s at | R49395 | 2881 | EST | 4.38 | g, | 0.00112 |
| rc R49476 at | R49476 | 2883 | EST | . 10.95 | ф | 0.00014 |
| rc_R52161_at | R52161 | 2893 | EST | 5.84 | g. | 0.03253 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change Direction | Direction | Pvalue |
|----------------|---------|--------|--|-----------------------|-----------|---------|
| rc R53109 f at | R53109 | 2899 | dimethylarginine dimethylaminohydrolase 2 | 3.31 | dn | 0.02389 |
| rc_R53109_r_at | R53109 | 2899 | dimethylarginine dimethylaminohydrolase 2 | 3.22 | dn | 0.00724 |
| rc_R54614 s at | R54614 | 2902 | EST | 3.24 | dn | 0.00526 |
| rc_R55470_at | R55470 | 2904 | EST | 3.59 | dn | 0.00515 |
| rc_R56095_s at | R56095 | 2906 | EST | 8.16 | dn | 0.00023 |
| R56678 at | R56678 | 2908 | EST | 3.81 | dn | 0.02242 |
| rc R56880 at | R56880 | 2909 | EST | 6.82 | dn | 0.02559 |
| rc_R58974_at | R58974 | 2910 | EST | 4.95 | dn | 0.00498 |
| rc_R59352_s_at | R59352 | 2915 | KIAA0296 gene product | 4.19 | dn | 0.00393 |
| rc R61297 s at | R61297 | 2920 | eukaryotic translation initiation factor 3, subunit 6 (48kD) | 6.42 | dn | 0.00126 |
| | R69700 | 2943 | EST | 6.71 | dn | 0.0021 |
| rc R70005 at | R70005 | 2944 | EST | 4.61 | dn | 0.00037 |
| rc_R70801_s_at | R70801 | 2950 | EST | 6.36 | dn | 0.00563 |
| rc_R71082_s_at | R71082 | 2951 | programmed cell death 5 | 3.6 | dn | 0.01338 |
| rc R71395 at | R71395 | 2952 | EST | 10.42 | dn | 0.00422 |
| rc_R73565_at | R73565 | 2959 | EST | 3.29 | dn | 0.03489 |
| l | | | O-linked N-acetylglucosamine (GlcNAc) transferase (UDP-N- | | | |
| | | | acetyigiucosamine:poiypeptide-N-acetyigiucosaminyi | | | |
| rc_R76782_s_at | R76782 | 2963 | transferase) | 5.83 | dn | 0.01126 |
| rc R77631 at | R77631 | 2967 | EST | 3.43 | dn | 0.00006 |
| rc_R79580_at | R79580 | 2970 | EST | 6.25 | dn | 0.00593 |
| rc_R87989_at | R87989 | 2979 | centrosome associated protein | 3.64 | dn | 0.00008 |
| rc_R91819_at | R91819 | 2984 | EST | 8.95 | dn | 0.0000 |
| rc R92994 s at | R92994 | 2990 | matrix metalloproteinase 12 (macrophage elastase) | 11.05 | 롸 | 0.00248 |
| rc R95966 i at | R95966 | 2997 | EST | 11.22 | dn | 0.00682 |
| rc R96924 s at | R96924 | 3001 | EST | 6.18 | dn | 0.03417 |
| rc R97759 at | R97759 | 3006 | serum/glucocorticoid regulated kinase | 5.99 | dn | 0.00221 |
| S54005 s at | S54005 | 3020 | thymosin, beta 10 | 7.03 | dn | 0.00334 |
| S56151 s at | S56151 | 3021 | milk fat globule-EGF factor 8 protein | 4.59 | dn | 0.0091 |
| S69272_s_at | S69272 | 3028 | protease inhibitor 6 (placental thrombin inhibitor) | 5.15 | dn | 0.00003 |
| | | | | | | |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affic ID | Genhank | Sea ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| | | | | | | |
| S73885 s at | S73885 | 3032 | transcription factor AP-4 (activating enhancer-binding protein 4) | 4.18 | dn | 0.00005 |
| S78187 at | S78187 | 3036 | cell division cycle 25B | 8.07 | dn | 60000.0 |
| S81914 at | S81914 | 3038 | immediate early response 3 | 5.46 | dn | 0.01798 |
| rc T03313 at | T03313 | 3042 | dyskeratosis congenita 1, dyskerin | 9.63 | dn | 0.00001 |
| 1 | T03438 | 3043 | EST | 8.18 | dn | 0.00032 |
| rc T03580 f at | T03580 | 3046 | pyruvate kinase, muscle | 24.91 | dn | 0.0001 |
| rc_T12599_f_at | T12599 | 3056 | ribosomal protein L21 | 3.54 | dn | 0.01437 |
| rc T15442 f at | T15442 | 3057 | calpain, large polypeptide L1 | 5.01 | dn | 0.00255 |
| rc T15473 at | T15473 | 3058 | muscle specific gene | 5.81 | dn | 0.02404 |
| rc T15477 at | T15477 | 3059 | EST | 3.61 | dn | 0.00005 |
| rc_T15903_s at | T15903 | 3063 | EST | 3.23 | dn | 0.01377 |
| rc_T16308_f_at | T16308 | 3069 | EST | 5.29 | dn | 0.00119 |
| rc T16983 s at | T16983 | 3074 | cleavage and polyadenylation specific factor 4, 30kD subunit | 5.23 | ф | 0.00075 |
| rc T23465 at | T23465 | 3081 | EST | 4.4 | dn | 0.017 |
| rc_T23490_i at | T23490 | 3082 | EST | 11.86 | d | 0.03242 |
| rc_T23516_f_at | T23516 | 3083 | 3-phosphoglycerate dehydrogenase | 5.38 | dn | 0.00001 |
| rc_T24068 s at | T24068 | 3088 | EST | 15.26 | dn | 0.00046 |
|) te | T25725 | 3091 | EST | 3.26 | dn | 66000.0 |
| rc_T26366_f at | T26366 | 3093 | EST | 30.43 | dn | 0.00153 |
| rc_T26471_at | T26471 | 3094 | EST | 4.62 | dn | 0.01091 |
| 1 | | | protein phosphatase 2 (formerly 2A), regulatory subunit A (PR | | | |
| rc T26513 at | T26513 | 3095 | 65), alpha isoform | 2.07 | dn | 0.0016 |
| rc T26574 s at | T26574 | 3096 | catenin (cadherin-associated protein), delta 1 | 3.17 | dn | 0.00828 |
| יי וי | T30193 | 3098 | protease, serine, 8 (prostasin) | 8.39 | dn | 0.00043 |
| rc T32072 s at | T32072 | 3102 | EST | 5.95 | đ | 0.00029 |
| rc T32108 at | T32108 | 3103 | EST | 96.9 | dn | 0.00723 |
| rc_T33489_s_at | T33489 | 3105 | EST | 8.04 | ф | 0.00469 |
| rc_T33508_s_at | T33508 | 3106 | phosphatidylinositol-4-phosphate 5-kinase, type II, beta | 3.31 | <u>а</u> | 0.00416 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | le Direction | Pvalue |
|----------------|---------|--------|--|-------------|--------------|---------|
| T35341 s at | T35341 | 3112 | EST | 4.73 | _ | 0.00057 |
| T35725 s at | T35725 | 3113 | EST | 3.4 | dn | 0.00149 |
| rc T40849 s at | T40849 | 3116 | maternal G10 transcript | 4.11 | dn | 0.00449 |
| rc_T47032_s_at | T47032 | 3124 | partner of RAC1 (arfaptin 2) | 3.27 | dn | 0.00503 |
| rc T47325 s at | T47325 | 3125 | EST | 5.63 | dn | 0.01015 |
| rc T47601 at | T47601 | 3126 | EST | 4.05 | dn | 0.00878 |
| 1 | | | ceroid-lipofuscinosis, neuronal 3, juvenile (Batten, Spielmeyer- | | | |
| rc_T47969_s_at | T47969 | 3128 | Vogt disease) | 3.03 | dn | 0.03452 |
| | | | eukaryotic translation initiation factor 3, subunit 3 (gamma, | | | |
| T48195 s at | T48195 | 3131 | 40kD) | 3.75 | dn | 0.00012 |
| rc T48293 f at | T48293 | 3133 | EST | 3.55 | dn | 0.01355 |
| rc T53404 at | T53404 | 3143 | EST | 10.68 | dn | 0.00582 |
| rc_T55004_s_at | T55004 | 3146 | EST | 4.83 | dn | 0.00156 |
| rc T55196 at | T55196 | 3147 | EST | 4.04 | dn | 0.00012 |
| rc_T58153_s_at | T58153 | 3154 | heat shock 105kD | 3.08 | dn | 0.01317 |
| rc_T58607_at | T58607 | 3155 | EST | 3.52 | dn | 0.04102 |
| rc_T59161_s_at | T59161 | 3159 | thymosin, beta 10 | 3.41 | dn | 0.01885 |
| rc T59668 s_at | T59668 | 3160 | lysyl oxidase | 3.28 | dn | 0.00588 |
| rc_T66935_at | T66935 | 3179 | EST | 3.97 | dn | 0.00188 |
| rc_T77733_s_at | T77733 | 3219 | tubulin, gamma 1 | 4.42 | ф | 0.00049 |
| rc T78922 s at | T78922 | 3222 | stem cell growth factor; lymphocyte secreted C-type lectin | 3.42 | g | 0.02419 |
| rc T91116 at | T91116 | 3252 | EST | 4.01 | dn | 0.02721 |
| rc_T92935_at | T92935 | 3255 | EST | 3.48 | dn | 0.03578 |
| rc_T95057_f at | T95057 | 3259 | EST | 10.39 | dn | 0.00003 |
| rc_T98284_at | T98284 | 3268 | EST | 4.47 | dn | 0.00054 |
| U01062_at | U01062 | 3273 | inositol 1,4,5-triphosphate receptor, type 3 | 7.41 | dn | 0 |
| U01147_at | U01147 | 3275 | active BCR-related gene | 3.22 | dn | 0.00103 |
| U02493_at | U02493 | 3279 | | 3.04 | dn | 0.0019 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affv ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|-------------------------------|------------------|--------------|--|----------------------|-----------|-------------------------------|
| | | | | c | 9 | 0 00085 |
| U03891_at U04313_at | U03891 U04313 | 3283 3284 | prorogin (similar to apolipoprotein b mixivo equaty protein) protease inhibitor 5 (maspin) | 4.54 | 하 와 | 0.02986 |
| I | | | | 6 | ; | 0 |
| U05875_at | U05875 | 3286 | interferon gamma receptor 2 (interferon gamma transducer 1) | 3.09 | d h | 0.00048 |
| U07969 s at | 007969 | 3289 | cadherin 17, Ll cadherin (liver-intestine) | 10.78 | dn | 0.02002 |
| U09117 at | U09117 | 3294 | phospholipase C, delta 1 | 2.96 | ф | 0.00001 |
| U09564_at | U09564 | 3295 | SFRS protein kinase 1 | 3.79 | dn | 0.00765 |
| U09770_at | 009770 | 3296 | cysteine-rich protein 1 (intestinal) | 13.03 | dn | 0.0072 |
| U11861_at | U11861 | 3298 | maternal G10 transcript | 3.8 | dn | 0.00001 |
| U12404 at | U12404 | 3299 | ribosomal protein L10a | 4.18 | dn | 0.00004 |
| U12465_at | U12465 | 3300 | ribosomal protein L35 | 4.69 | dn | 0.00001 |
| U14968 at | U14968 | 3303 | ribosomal protein L27a | 4.01 | dn | 0.00003 |
| U14969 at | U14969 | 3304 | ribosomal protein L28 | 4.63 | dn | 0.00004 |
| U14970_at | U14970 | 3305 | ribosomal protein S5 | 3.45 | dn | 0.00915 |
| U14971 at | U14971 | 3306 | ribosomal protein S9 | 3.93 | dn | 0.00026 |
| U14972 at | U14972 | 3307 | ribosomal protein S10 | 5.24 | dn | 0.00077 |
| U14973_at | U14973 | 3308 | ribosomal protein S29 | 3.1 | dn | 0.00028 |
| 1145008 at | 115008 | 3309 | small nuclear ribonucleoprotein D2 polypeptide (16.5kD) | 4.9 | g | 0.00396 |
| U17077_at | U17077 | 3314 | BENE protein | 4.98 | ф | 0.00366 |
| | U17760 | 3315 | laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 (125kD)) | 3.54 | dn | 0.01853 |
| U20499_at | U20499 | 3321 | sulfotransferase family 1A, phenol-preferring, member 3 | 5.5 | dn | 0.00299 |
| U20758_rna1_at | U20758 | 3323 | secreted phosphoprotein 1 (osteopontin, borle statoprotein 1, early T-lymphocyte activation 1) | 6.12 | 윰 | 0.03448 |
| U21049_at U22376_cds2_s_at | U21049 U22376 | 3325 | epimelial protein up-regulated in calcinolità, inellibrate associated protein 17 v-myb avian myeloblastosis viral oncogene homolog | 7.53 3.34 4.37 | 요요 | 0.01667 0.03416 0.00045 |
| U25/89_at | 69/670 | 4000 | IIDOSOINAI PLOTEILI EZ I | è | <u>}</u> | |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change Direction | Direction | Pvalue |
|------------------|----------|--------|---|-----------------------|----------------|---------|
| U26726_at | U26726 | 3336 | hydroxysteroid (11-beta) dehydrogenase 2 | 3.45 | dn | 0.02342 |
| | • | | fucosyltransferase 3 (galactoside 3(4)-L-fucosyltransferase, | | | |
| U27328_s_at | U27328 | 3338 | Lewis blood group included) | 3.05 | an | 0.03224 |
| U31556_at | U31556 | 3346 | E2F transcription factor 5, p130-binding | 4.14 | - 9 | 0.01157 |
| U33286_at | U33286 | 3349 | chromosome segregation 1 (yeast homolog)-like | 3.39 | d an | 0.00939 |
| | | | solute carrier family 6 (neurotransmitter transporter, creatine), | | • | |
| U36341_rna1_at | U36341 | 3351 | member 8 | 3.05 | an | 0.02622 |
| U37689_at | U37689 | 3356 | polymerase (RNA) II (DNA directed) polypeptide H | 3.65 | dn | 0.00044 |
| | | | potassium voltage-gated channel, KQT-like subfamily, member | | - | |
| U40990_at | U40990 | 3359 | _ | 3.18 | an | 0.00093 |
| U42408_at | U42408 | 3361 | ladinin 1 | 6.8 | gn | 0.00305 |
| U43901_rna1_s_at | U43901 | 3362 | laminin receptor 1 (67kD, ribosomal protein SA) | 3.11 | an | 0.03145 |
| U46692_rna1_at | U46692 | 3367 | cystatin B (stefin B) | 5.54 | . an | 0.00016 |
| U47025_s_at | U47025 | 3368 | phosphorylase, glycogen; brain | 8.52 | . an | 0.00134 |
| U48705_rna1_s_at | U48705 | 3370 | discoidin domain receptor family, member 1 | 5.94 | . an | 0.01323 |
| U51095_at | U51095 | 3382 | caudal type homeo box transcription factor 1 | 4.76 | g. | 0.02664 |
| U51478_at | . U51478 | 3385 | ATPase, Na+/K+ transporting, beta 3 polypeptide | 5.75 | g | 0.00007 |
| | | | solute carrier family 1 (neutral amino acid transporter), member | | • | |
| U53347_at | U53347 | 3389 | ro. | 3.81 | dn | 0.00273 |
| U53830_at | U53830 | 3391 | interferon regulatory factor 7 | 3.94 | . an | 0.03147 |
| U58682_at | U58682 | 3396 | ribosomal protein S28 | 6.09 | . <u>a</u> | 0.00001 |
| U62392_at | U62392 | 3403 | zinc finger protein 193 | 3.18 | - ф | 0.00269 |
| U62962_at | U62962 | 3404 | eukaryotic translation initiation factor 3, subunit 6 (48kD) | 4.05 | an | 0.0047 |
| U67171_at | U67171 | 3409 | selenoprotein W, 1 | 3.08 | Ωn | 0.0047 |
| U73379_at | U73379 | 3418 | ubiquitin carrier protein E2-C | 8.32 | - an | 0.00101 |
| U73843_at | U73843 | 3421 | E74-like factor 3 (ets domain transcription factor) | 5.75 | an | 0.00017 |
| U75285_rna1_at | U75285 | 3422 | apoptosis inhibitor 4 (survivin) | 4.46 | . an | 0.02212 |
| U76366_s_at | U76366 | 3424 | Treacher Collins-Franceschetti syndrome 1 | 3.44 | . <u>a</u> | 0.00021 |
| U78027_rna3_at | U78027 | 3429 | EST | 4.15 | 9 | 0.00295 |
| U78095_at | U78095 | 3430 | serine protease inhibitor, Kunitz type, 2 | 18.85 | - <u>a</u> | 0 |
| | | | | | - | |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | e Direction | Pvalue |
|----------------|----------|--------|--|-------------|-------------|---------|
| | | | | | | |
| U78525 at | U78525 | 3432 | eukaryotic translation initiation factor 3, subunit 9 (eta, 116kD) | 4.68 | dn | 0.00132 |
| U78556 at | U78556 | 3433 | cisplatin resistance associated | 5.77 | dn | 0.00241 |
| U79266_at | U79266 | 3434 | protein predicted by clone 23627 | 3.49 | dn | 0.00004 |
| U79725_at | U79725 | 3438 | glycoprotein A33 (transmembrane) | 8.57 | dn | 0.00299 |
| U83246_at | U83246 | 3443 | copine I | 3.57 | dn | 0.01672 |
| U84720 at | U84720 | 3446 | RAE1 (RNA export 1, S.pombe) homolog | 3.37 | dn | 0.03586 |
| U85773 at | · U85773 | 3449 | phosphomannomutase 2 | 3.94 | dn | 0.00288 |
| U86409_at | U86409 | 3450 | EST | 3.38 | dn | 0.00003 |
| U89606_at | U89606 | 3452 | pyridoxal (pyridoxine, vitamin B6) kinase | 3.58 | 함 | 0.00322 |
| U90549 at | U90549 | 3456 | high-mobility group (nonhistone chromosomal) protein 17-like 3 | 3.2 | dn | 0.0401 |
| U90913_at | U90913 | 3459 | Tax interaction protein 1 | 4.35 | dn | 0.00159 |
| U93205_at | U93205 | 3461 | chloride intracellular channel 1 | 6.14 | dn | 0.00058 |
| U93868 at | U93868 | 3463 | polymerase (RNA) III (DNA directed) (32kD) | 3.5 | dn | 0.01235 |
| rc W02041 at | W02041 | 3466 | EST | 4.83 | ф | 0.00158 |
| rc_W20391 s at | W20391 | 3479 | kinesin-like 2 | 3.98 | d d | 0.01788 |
| W28362 at | W28362 | 3488 | KIAA0974 protein | 3.98 | dn | 0.00626 |
| rc_W31382_at | W31382 | 3495 | EST | 4 | dn | 0.00058 |
| rc_W37680_at | W37680 | 3503 | EST | 3.55 | dn | 0.01036 |
| rc_W37937_at | W37937 | | EST | 3.07 | ф | 0.00776 |
| | | | myeloid/lymphoid or mixed-lineage leukemia (trithorax | | | |
| rc W38044 s at | W38044 | | (Drosophila) homolog); translocated to, 7 | 7.28 | d | 0.03105 |
| W39183 s at | . W39183 | 3508 | KIAA0601 protein | 3.66 | 롸 | 0.00018 |
| rc W42627 f at | W42627 | 3511 | EST | 3.56 | dn | 0.00198 |
| rc_W42957_at | W42957 | 3516 | calmodulin 2 (phosphorylase kinase, delta) | 15.22 | dn | 0.00007 |
| rc_W44557_at | W44557 | 3518 | chromosome 1 open reading frame 2 | 4.32 | dn | 0.00128 |
| rc_W44733_at | W44733 | 3519 | EST | 3.02 | ф | 0.00097 |
| rc_W45487_s_at | W45487 | 3524 | dynamin 2 | 4.2 | dn | 0.00325 |
| rc_W49574_at | W49574 | 3538 | EST | 5.97 | dn | 0.00045 |
| rc_W49661_s_at | W49661 | 3539 | FK506-binding protein 9 (63 kD) | 3.01 | dn | 0.02259 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | ge Direction | Pvalue |
|----------------|---------|--------|--|-------------|----------------|---------|
| W52858 at | W52858 | 3545 | DKFZP564F0522 protein | 4.38 | dп | 0.00088 |
| rc_W60486_at | W60486 | 3558 | | 4.29 | - c | 0.00964 |
| rc_W67251_s_at | W67251 | 3570 | EST | 6.13 | d | 0.01463 |
| rc_W70336_at | W70336 | 3579 | EST | 3.65 | g | 0.01776 |
| rc_W72276_at | W72276 | 3583 | EST | 27.39 | dn | 0.00183 |
| | | | protein tyrosine phosphatase, receptor type, f polypeptide | | | |
| rc_W72861_at | W72861 | 3586 | (PTPRF), interacting protein (liprin), alpha 3 | 3.01 | dn | 0.00055 |
| rc_W73189_at | W73189 | 3589 | EphB2 | 3.69 | d. | 0.02909 |
| rc_W73914_at | W73914 | 3595 | EST | 3.5 | 음 | 0.04782 |
| rc_W74233_s_at | W74233 | 3597 | related RAS viral (r-ras) oncogene homolog | 6.02 | dn | 0.01815 |
| W76097_at | W76097 | 3599 | EST | 3.83 | dn | 0.00001 |
| rc_W78057_at | W78057 | 3600 | EST | 90.6 | dn | 0.0034 |
| rc_W80730_at | W80730 | 3607 | EST | 6.59 | d | 0.00425 |
| rc_W80763_at | W80763 | 3608 | EST | 3.83 | dn | 0.01319 |
| rc_W81375_at | W81375 | 3613 | EST | 3.58 | dn | 0.00322 |
| rc_W81540_at | W81540 | 3614 | serine/threonine kinase 24 (Ste20, yeast homolog) | 6.72 | dn | 0.00164 |
| rc_W90146_f_at | W90146 | 3644 | EST | 6.23 | dn | 0.01558 |
| rc_W92207_at | W92207 | 3651 | EST | 6.77 | dn | 0.00002 |
| rc_W92449_at | W92449 | 3652 | EST | 31.67 | dn | 0.00011 |
| rc_W92608_s_at | W92608 | 3653 | BAI1-associated protein 3 | 5.12 | dn | 0.00075 |
| rc_W93726_s_at | W93726 | 3656 | protease inhibitor 5 (maspin) | 16.48 | ď | 0.00014 |
| rc_W93943_at | W93943 | 3657 | EST | 4.3 | dn | 0.00296 |
| W95348_at | W95348 | 3663 | HSPC113 protein | 10.89 | dn | 0.01065 |
| rc_W95477_at | W95477 | 3664 | EST | 26.51 | dn | 0.00161 |
| X03342_at | X03342 | 3675 | ribosomal protein L32 | 4.09 | ď | 0.00008 |
| AFFX-CreX-5_at | X03453 | 3677 | EST | 3.03 | dn | 0.0025 |
| X04347_s_at | X04347 | 3680 | heterogeneous nuclear ribonucleoprotein A1 | 7.26 | dn | 0.00018 |
| X05610_at | X05610 | 3685 | collagen, type IV, alpha 2 | 3.58 | dn | 0.01351 |
| X06617_at | X06617 | 3687 | ribosomal protein S11 | 4.32 | đ | 0.0002 |
| X07820_at | X07820 | 3695 | matrix metalloproteinase 10 (stromelysin 2) | 3.49 | dn | 0.00689 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|-----------|---------|
| X12447 at | X12447 | 3696 | aldolase A, fructose-bisphosphate | 10.52 | dn | 0.00038 |
| X13956 at | X13956 | 3701 | EST | 3.2 | dn | 0.00321 |
| X14850 at | X14850 | 3706 | H2A histone family, member X | 4.11 | dn | 0.0001 |
| X15940_at | X15940 | 3709 | ribosomal protein L31 | 4.45 | dn | 0.00004 |
| X17093 at | X17093 | 3716 | EST | 4.82 | dn | 0.00176 |
| X17206 at | X17206 | 3718 | ribosomal protein S2 | 5.14 | dn | 0 |
| X17567 s at | X17567 | 3719 | small nuclear ribonucleoprotein polypeptides B and B1 | 3.6 | dn | 0.00586 |
| X51466 at | X51466 | 3720 | eukaryotic translation elongation factor 2 | 3.25 | dn | 0.00019 |
| X51521_at | X51521 | 3721 | villin 2 (ezrin) | 3.89 | dn | 0.00001 |
| X52851 rna1 at | X52851 | 3725 | EST | 3.38 | dn | 0.0001 |
| X52966 at | X52966 | 3726 | ribosomal protein L35a | 3.93 | dn | 0.00217 |
| X53331 at | X53331 | 3727 | matrix Gla protein | 3.66 | dn | 0.04038 |
| X54667 s at | X54667 | 3731 | cystatin S, cystatin SN | 8.53 | dn | 0.00059 |
| X55715 at | X55715 | 3735 | ribosomal protein S3 | 3.72 | dn | 0.00755 |
| X55954 at | X55954 | 3736 | ribosomal protein L23 | 3.81 | dn | 0.00025 |
| X56494 at | X56494 | 3738 | pyruvate kinase, muscle | 22.97 | dn | 0.00001 |
| X56932_at | X56932 | 3740 | ribosomal protein L13a | 3.26 | dn | 0 |
| X56997 rna1 at | X56997 | 3741 | ubiquitin A-52 residue ribosomal protein fusion product 1 | 3.18 | d | 0.00006 |
| X57348 s at | X57348 | 3744 | stratifin | 12.53 | dn | 0.0013 |
| X62535 at | X62535 | 3756 | diacylglycerol kinase, alpha (80kD) | 4.72 | dn | 0.00315 |
| X62691_at | X62691 | 3757 | ribosomal protein S15a | 4.09 | dn | 0.00005 |
| X63527_at | X63527 | 3761 | ribosomal protein L19 | 3.17 | dn | 0.02488 |
| X63629_at | X63629 | 3762 | cadherin 3, P-cadherin (placental) | 3.02 | ф | 0.01654 |
| X64364_at | X64364 | 3764 | basigin | 6.45 | dn | 0.00041 |
| X64707 at | X64707 | 3765 | ribosomal protein L13 | 4.28 | dn | 0.00257 |
| X65614_at | X65614 | 3767 | S100 calcium-binding protein P | 12.2 | ф | 0.00065 |
| X66364_at | X66364 | 3770 | cyclin-dependent kinase 5 | 3.55 | dn | 0.02824 |
| | | | proteasome (prosome, macropain) subunit, beta type, 9 (large | Ċ | <u>!</u> | 2007 |
| X66401_cds1_at | X66401 | 3771 | _ | 3.68 | dn ! | 0.01383 |
| X66899_at | X66899 | 3772 | Ewing sarcoma breakpoint region 1 | 4.72 | 읔 | 0.00011 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | e Direction | Pvalue |
|------------------|---------|--------|---|-------------|----------------|---------|
| X67247 rna1 at | X67247 | 3774 | mitogen-activated protein kinase kinase kinase 8 | 3.48 | dn | 0.00012 |
| X67325 at | X67325 | 3775 | interferon, alpha-inducible protein 27 | 9.67 | dn | 0.03245 |
| X68314_at | X68314 | 3778 | glutathione peroxidase 2 (gastrointestinal) | 14.4 | g | 0.00222 |
| X68688 rna1 s at | X68688 | 3780 | zinc finger protein 33a (KOX 31) | 4.7 | g. | 0.00062 |
| | X68688 | 3780 | zinc finger protein 33a (KOX 31) | 3.62 | dn | 0.00319 |
| | X69150 | 3782 | ribosomal protein S18 | 4.5 | 슠 | 0.00003 |
| X69391_at | X69391 | 3783 | ribosomal protein L6 | 3.18 | 9 | 0.00003 |
| i | | | CD47 antigen (Rh-related antigen, integrin-associated signal | | | |
| X69398 at | X69398 | 3784 | fransducer) | 3.68 | d | 0.02332 |
| X69654_at | X69654 | 3785 | ribosomal protein S26 | 3.11 | dn | 0.02683 |
| I | | | ATP synthase, H+ transporting, mitochondrial F0 complex, | | | |
| X69908_rna1_at | 80669X | 3786 | subunit c (subunit 9), isoform 2 | 3.73 | dn | 0.00685 |
| | | | macrophage stimulating 1 receptor (c-met-related tyrosine | | | |
| X70040 at | X70040 | 3788 | kinase) | 4 | d _n | 0.00402 |
| X74929 s at | X74929 | 3792 | keratin 8 | 4.73 | dn | 0.00018 |
| X76180_at | X76180 | 3795 | sodium channel, nonvoltage-gated 1 alpha | 11.68 | dn | 0 |
| X77588 s at | X77588 | 3799 | N-acetyltransferase, homolog of S. cerevisiae ARD1 | 3.91 | ф | 0.0221 |
| X78687_at | X78687 | 3800 | sialidase 1 (lysosomal sialidase) | 3.18 | ф | 0.031 |
| X79234 at | X79234 | 3803 | ribosomal protein L11 | 3.59 | ф | 0.00051 |
| X79882_at | X79882 | 3805 | lung resistance-related protein | 6.71 | dn | 0.0362 |
| X80198 at | X80198 | 3807 | steroidogenic acute regulatory protein related | 3.03 | dn | 0.00044 |
| X80822_at | X80822 | 3808 | ribosomal protein L18a | 4.13 | dn | 0.0006 |
| X80822_f_at | X80822 | 3808 | ribosomal protein L18a | 3.08 | ф | 0.02481 |
| X80909 at | 60608X | 3809 | nascent-polypeptide-associated complex alpha polypeptide | 3.84 | dn | 0.00399 |
| X83228 at | X83228 | 3810 | cadherin 17, Ll cadherin (liver-intestine) | 10.58 | dn | 0.02147 |
| X89960_at | 09668X | 3818 | EST | 9.87 | dn | 0 |
| | | | high-mobility group (nonhistone chromosomal) protein isoform I- | | | |
| X92518_s_at | X92518 | 3825 | | 3.12 | d | 0.00638 |
| X93036_at | X93036 | 3830 | FXYD domain-containing ion transport regulator 3 | 42.36 | d | 0.00167 |

Table 6A. Genes up regulated in metastatic tissue versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| X95404 at | X95404 | 3834 | cofilin 1 (non-muscle) | 3.18 | dn | 0.00104 |
| X98482 r at | X98482 | 3841 | EST | 5.03 | dn | 0.00002 |
| X99133_at | X99133 | 3842 | lipocalin 2 (oncogene 24p3) | 6.27 | dn | 0.0453 |
| Y00503 at | Y00503 | 3849 | keratin 19 | 14.19 | dn | 0.00217 |
| Y00705 at | Y00705 | 3850 | serine protease inhibitor, Kazal type 1 | 9.83 | dn | 0.03697 |
| Y10807 s at | Y10807 | 3860 | HMT1 (hnRNP methyltransferase, S. cerevisiae)-like 2 | 4.28 | dn | 0.00124 |
| Z23090 at | Z23090 | 3868 | heat shock 27kD protein 1 | 7.67 | dn | 0.00008 |
| Z24727 at | Z24727 | 3871 | tropomyosin 1 (alpha) | 4.47 | dn | 0.00121 |
| ı | | | nuclear factor of kappa light polypeptide gene enhancer in B- | | | |
| Z25749 ma1 at | Z25749 | 3872 | cells inhibitor-like 2 | 4.41 | dn | 0.00031 |
| Z26876 at | Z26876 | 3874 | ribosomal protein L38 | 4.43 | d | 0.00022 |
| Z28407 at | Z28407 | 3876 | ribosomal protein L8 | 6.53 | dn | 0.00004 |
| Z30643 at | Z30643 | 3879 | chloride channel Ka | 3.86 | dn | 0.00204 |
| rc Z38150 s at | Z38150 | 3883 | EST | 3.06 | dn | 0.00049 |
| rc_Z38266_at | Z38266 | 3886 | EST | 6.58 | dn | 0.01909 |
| rc Z38729 at | Z38729 | 3894 | EST | 3.13 | ф | 0.04514 |
| rc_Z38909_at | Z38909 | 3897 | EST | 3.55 | dn | 0.03195 |
| rc_Z39079_at | Z39079 | 3900 | KIAA1058 protein | 4.08 | dn | 0.01781 |
| rc_Z39191_at | Z39191 | 3901 | EST | 8.84 | dn | 0.00011 |
| rc_Z39200_at | Z39200 | 3902 | EST | 3.29 | dn | 0.00586 |
| rc_Z39429_at | Z39429 | 3906 | EST | 7.84 | dn | 0.00045 |
| rc Z39930 f at | Z39930 | 3914 | EST | 3.07 | dn | 0.00002 |
| rc Z40583 f at | Z40583 | 3922 | EST | 4.24 | ф | 0.02375 |
| rc Z40898_at | Z40898 | 3925 | EST | 3.71 | dn | 0.01168 |
| rc_Z40945_at | Z40945 | 3927 | trinucleotide repeat containing 15 | 7.82 | dn | 0.00007 |
| rc Z41103 at | Z41103 | 3929 | trinucleotide repeat containing 15 | 3.83 | dn | 0.00444 |
| rc_Z41740_s_at | Z41740 | 3935 | EST | 9.76 | dn | 0.00049 |
| rc_Z41798_s_at | Z41798 | 3937 | EST | 9 | ф | 0.00073 |
| Z68228 s at | Z68228 | 3947 | junction plakoglobin | 3.3 | dn | 0.0237 |
| Z74615_at | Z74615 | 3949 | collagen, type I, alpha 1 | 10.47 | dn | 0.00064 |
| Z74616_s_at | Z74616 | 3950 | collagen, type I, alpha 2 | 4.83 | dn | 0.02364 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| rc_AA001603_at | AA001603 | က | EST | 4.31 | down | 0.00883 |
| rc_AA001604_at | AA001604 | 4 | EST | 3.33 | down | 0.0215 |
| AA004231_at | AA004231 | 7 | EST | 3.16 | down | 0.03067 |
| rc_AA004521_at | AA004521 | 80 | prostate cancer overexpressed gene 1 | 8.03 | down | 0.00027 |
| rc_AA004905_at | AA004905 | 7 | KIAA0937 protein | 4.63 | down | 0.00082 |
| rc_AA005202_at | AA005202 | 12 | retinol-binding protein 4, interstitial | 3.18 | down | 0.00106 |
| rc_AA005358_at | AA005358 | 4 | EST | 5.2 | down | 0.00318 |
| rc_AA007629_at | AA007629 | 19 | EST | 8.01 | down | 0.00001 |
| rc_AA009719_at | AA009719 | 20 | peroxisomal membrane protein 2 (22kD) | 47.12 | down | 0.00008 |
| rc_AA010205_at | AA010205 | 23 | EST | 7.41 | down | 0 |
| rc_AA010360_at | AA010360 | 54 | EST | 6.55 | down | 0.00027 |
| rc_AA010530_at | AA010530 | 25 | EST | 3.74 | down | 0.0481 |
| rc_AA010619_at | AA010619 | 27 | EST | 8.55 | down | 0.00057 |
| AA010750_at | AA010750 | 28 | calmodulin 1 (phosphorylase kinase, delta) | 10.22 | down | 0.00959 |
| rc_AA015768_at | AA015768 | 34 | EST | 15.3 | down | 0.00008 |
| rc_AA017146_at | AA017146 | 36 | EST | 10.1 | down | 0.00052 |
| rc_AA017192_at | AA017192 | 37 | EST | 3.43 | down | 0.04865 |
| rc_AA018867_at | AA018867 | 33 | EST | 42.87 | down | 0.00002 |
| rc_AA021623_s_at | AA021623 | 43 | insulin induced gene 1 | 11.71 | down | 0.00094 |
| rc_AA025930_at | AA025930 | 25 | EST | 3.59 | down | 0.00372 |
| rc_AA031543_s_at | AA031543 | 89 | translocation protein 1 | 5.92 | down | 0.00405 |
| AA031548 at | AA031548 | 69 | cell division cycle 42 (GTP-binding protein, 25kD) | 3.65 | down | 0.03029 |
| rc_AA032005_at | AA032005 | 71 | EST | 5.3 | down | 0.01202 |
| AA032048_at | AA032048 | 72 | EST | 5.45 | down | 0.00383 |
| rc_AA032250_at | AA032250 | 73 | EST | 3.56 | down | 0.0009 |
| rc_AA034030_at | AA034030 | 75 | methylmalonyl Coenzyme A mutase | 14.32 | down | 0.00004 |
| rc_AA035245_s_at | AA035245 | 79 | aldehyde oxidase 1 | 69.82 | down | 0.00117 |
| rc_AA035457_at | AA035457 | 80 | EST | 10.06 | down | 0.00085 |
| rc_AA035638_at | AA035638 | 82 | EST | 9.91 | down | 0.00541 |
| rc_AA036662_s_at | AA036662 | 83 | EST | 4.16 | down | 0.00235 |
| rc_AA037357_f_at | AA037357 | 82 | EST | 3.53 | down | 0.02129 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| rc AA039335 s at | AA039335 | 88 | coagulation factor XII (Hageman factor) | 32 | down | 0.0029 |
| rc AA039616 at | AA039616 | 90 | EST | 9.36 | down | 60000.0 |
| AA039806 at | AA039806 | 91 | msh (Drosophila) homeo box homolog 1 | 3.53 | down | 0.00114 |
| rc_AA040087_at | AA040087 | 92 | EST | 4.13 | down | 0.00123 |
| | AA040270 | 93 | EST | 3.03 | down | 0.01367 |
| rc_AA040291_at | AA040291 | 94 | KIAA0669 gene product | 3.55 | down | 0.00308 |
| rc_AA043501_at | AA043501 | 86 | v-maf musculoaponeurotic fibrosarcoma (avian) | 3.81 | down | 0.01304 |
| AA044622 at | AA044622 | 103 | EST | 3.62 | down | 0.03789 |
| AA044755 s at | AA044755 | 104 | EST | 6.7 | down | 0.01228 |
| AA044842_at | AA044842 | 105 | Autosomal Highly Conserved Protein | 5.21 | down | 6000.0 |
| AA045870_at | AA045870 | 108 | EST | 5.93 | down | 0.00017 |
| rc AA046457 at | AA046457 | 1 | EST | 3.2 | down | 0.00513 |
| AA046674 at | AA046674 | 112 | EST | 5.17 | down | 0.02561 |
| rc_AA046747_at | AA046747 | 114 | EST | 4.82 | down | 0.00022 |
| AA046840 at | AA046840 | 115 | CCAAT/enhancer binding protein (C/EBP), delta | 3.79 | down | 0.03319 |
| AA047151_at | AA047151 | 116 | EST | 7.13 | down | 0.00007 |
| rc_AA047187_at | AA047187 | 117 | EST | 3.04 | down | 0.04306 |
| rc_AA047290_at | AA047290 | 118 | EST | 3.39 | down | 0.00024 |
| rc_AA052980_at | AA052980 | 122 | EST | 4.52 | down | 0.023 |
| rc_AA055992_at | AA055992 | 136 | calumenin | 3.51 | down | 0.00604 |
| AA056170_at | AA056170 | 137 | EST | 3.82 | down | 0.0083 |
| rc_AA056247_at | AA056247 | 138 | EST | 3.48 | down | 0.03277 |
| rc_AA056482_at | AA056482 | 141 | EST | 4.82 | down | 0.00199 |
| AA057678 | AA057678 | 143 | EST | 6.88 | down | 0.00078 |
| ĀA059489 at | AA059489 | 145 | RGC32 protein | 3.74 | down | 0.00734 |
| rc AA062744 at | AA062744 | 147 | EST | 3.31 | down | 0.01909 |
| rc_AA065173_at | AA065173 | 148 | EST | 4.08 | down | 0.00377 |
| rc_AA069456 at | AA069456 | 149 | KIAA0438 gene product | 3.47 | down | 0.02718 |
| rc AA069768 s at | AA069768 | 151 | hevin | 4.62 | down | 0.00202 |
| AA070090_at | AA070090 | 152 | EST | 3.24 | down | 0.00804 |
| rc_AA070091_at | AA070091 | 153 | EST | 5 | down | 0 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Pvalue | 0.00786 | 0.01897 | 0.00395 | 0.00029 | 0.00616 | | 0.00359 | 0.00167 | 0.00864 | 0.0419 | 0.00004 | 0.00012 | 0.02543 | 0.02072 | 0.01704 | 0.00501 | 0.02532 | 0.00009 | 0.03924 | 0.04783 | 0.0257 | 0.00062 | 0.00003 | 0.04069 | 0.0071 | 0.00822 | 0.00023 | 0.00004 | 0.00084 | 0.01028 |
|-----------------|--|----------------|----------------|----------------|------------------------------|--|--------------------------------|----------------------|----------------|------------------|---------------------------|----------------|----------------|---------------------------------------|------------------------------------|----------------------|------------------------------|-------------------------------|-------------|---------------|-------------------------|----------------|----------------------------------|------------------------------|------------------|----------------|----------------|------------------|--|----------------------------|
| Direction | down | down | down | down | down | | down | down | down | down | down | down | down | down | down | down | down | down | down | down | down | down | down | down | down | down | down | down | down | down |
| Fold Change | 11.05 | 3.12 | 3.23 | 3.78 | 10.88 | | 7.83 | 9.11 | 4.05 | 3.19 | 40.87 | 5.8 | 3.23 | 11.72 | 4.61 | 5.58 | 3.46 | 13.97 | 3.82 | 3.52 | 5.12 | 7.33 | 9.07 | 3.7 | 3.14 | 8.46 | 4.19 | 8.5 | 3.37 | 5.12 |
| Known Gene Name | macrophage receptor with collagenous structure | EST | EST | EST | SEC14 (S. cerevisiae)-like 2 | solute carrier family 25 (mitochondrial carrier; | citrate transporter), member 1 | DKFZP566F123 protein | EST | ubiquitin-like 3 | UDP glycosyltransferase 1 | EST | EST | superoxide dismutase 2, mitochondrial | diaphanous (Drosophila, homolog) 1 | ribosomal protein S6 | bone morphogenetic protein 6 | HLA-B associated transcript-3 | EST | EST | zinc finger protein 216 | EST | myosin, light polypeptide kinase | GDP dissociation inhibitor 2 | leptin receptor | EST | EST | EST | acyl-Coenzyme A dehydrogenase, long chain UDP-N-acteylglucosamine pyrophosphorylase 1; | Sperm associated antigen 2 |
| Sed ID | 161 | 162 | 168 | 169 | 170 | | 174 | 175 | 179 | 180 | 183 | 185 | 188 | 190 | 191 | 192 | 197 | 198 | 200 | 201 | 204 | 206 | 207 | 210 | 213 | 214 | 217 | 222 | 223 | 228 |
| Genbank | AA074885 | AA074891 | AA076238 | AA076249 | AA076326 | | AA079758 | AA083812 | AA084408 | AA084668 | AA085987 | AA086201 | AA088698 | AA090257 | AA090434 | AA090439 | AA092596 | AA092716 | AA093923 | AA094507 | AA094999 | AA099225 | AA099391 | AA099589 | AA101055 | AA101235 | AA101632 | AA112101 | AA112209 | AA114949 |
| Affy ID | rc AA074885 at | rc_AA074891_at | rc_AA076238_at | rc_AA076249_at | rc_AA076326_at | ! | rc_AA079758_f_at | rc_AA083812_at | rc_AA084408_at | rc_aa084668_at | rc AA085987_s_at | rc_AA086201_at | rc_AA088698_at | AA090257_at | AA090434_at | AA090439_at | AA092596_at | AA092716_at | AA093923_at | AA094507_s_at | AA094999_at | rc_AA099225_at | AA099391_s_at | rc_AA099589_s_at | rc_AA101055_s_at | rc_AA101235_at | rc_AA101632_at | rc_AA112101_f_at | ω, | AA114949_at |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| rc AA121140 at | AA121140 | 235 | EST | 3.33 | down | 0.00058 |
| rc AA122345 f at | AA122345 | 238 | glutamate dehydrogenase 1 | 16.53 | down | 0.00004 |
| rc_AA125831_r_at | AA125831 | 241 | myosin, light polypeptide kinase | 8.18 | down | 0.00039 |
| rc AA125856 at | AA125856 | 242 | EST | 3.17 | down | 0.01545 |
| rc_AA125861_at | AA125861 | 243 | EST | 3.69 | down | 0.01547 |
| rc_AA126059_at | AA126059 | 246 | EST | 3.08 | down | 0.00706 |
| rc AA126722 s at | AA126722 | 251 | O-6-methylguanine-DNA methyltransferase | 8.26 | down | 0.002 |
| rc_AA127444_at | AA127444 | 252 | EST | 3.56 | down | 0.0291 |
| rc_AA127514_at | AA127514 | 253 | EST | 3.4 | down | 0.00045 |
| rc_AA133215_at | AA133215 | 277 | calcitonin receptor-like receptor activity modifying | 4.55 | down | 0.02092 |
| rc_AA133296_at | AA133296 | 278 | EST | 4.23 | down | 0.00041 |
| rc_AA133439_at | AA133439 | 279 | EST | 4.07 | down | 0.00022 |
| rc_AA134549_at | AA134549 | 288 | EST | 3.36 | down | 0.03438 |
| rc AA135558 s at | AA135558 | 293 | peptidase D | 4.17 | down | 0.0068 |
| rc AA135958_at | AA135958 | 296 | EST | 4.02 | down | 0.00012 |
| rc_AA136079_at | AA136079 | 297 | EST | 4.26 | down | 0.0057 |
| rc AA142849 at | AA142849 | 306 | EST | 7.59 | down | 0.00804 |
| AA143019_at | AA143019 | 309 | EST | 6.75 | down | 0.00109 |
| rc_aa147646_s_at | AA147646 | 317 | DKFZP586A0522 protein | 21.82 | down | 0 |
| rc_AA148480_s_at | AA148480 | 318 | flavin containing monooxygenase 5 | 19.64 | down | 0 |
| rc_AA148923_at | AA148923 | 321 | decidual protein induced by progesterone | 13.2 | down | 0.00257 |
| rc_AA149253_at | AA149253 | 323 | EST | 5.12 | down | 0.00863 |
| rc_AA150776_at | AA150776 | 330 | EST | 10.45 | down | 0.00015 |
| rc_AA150891_at | AA150891 | 331 | EST | 4.22 | down | 0.01692 |
| rc AA151210 at | AA151210 | 333 | EST | 4.61 | down | 0.00008 |
| rc_AA151676_at | AA151676 | 337 | peptidyl arginine deiminase, type II | 4.01 | down | 0.00911 |
| rc_AA156336_at | AA156336 | 341 | nuclear receptor co-repressor 1 | 3.69 | down | 0.01276 |
| l 1 | | | 4-nitrophenylphosphatase domain and non- | | | |
| rc_AA156565_at | AA156565 | 344 | neuronal SNAP25-like 1 | 15.01 | down | 0.01387 |
| rc_AA157112_at | AA157112 | 345 | EST | 3.94 | down | 0.02571 |
| AA157520_at | AA157520 | 347 | EST | 3.18 | down | 0.00516 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| | | | aldo-keto reductase family 7, member A2 | | | |
| rc_AA157799_at | AA157799 | 348 | (aflatoxin aldehyde reductase) | 5.96 | down | 0 |
| rc_AA164586_s_at | AA164586 | 326 | estrogen receptor 1 | 5.8 | down | 0.00182 |
| rc_AA167565_at | AA167565 | 362 | EST | 8.08 | down | 0.00046 |
| rc_AA171694_at | AA171694 | 366 | ceruloplasmin (ferroxidase) | 21.23 | down | 0.00179 |
| rc_AA172372_at | AA172372 | 370 | EST | 6.48 | down | 0.00344 |
| rc_AA176233_at | AA176233 | 376 | EST | 11.44 | down | 0.01856 |
| rc_AA179004_at | AA179004 | 377 | EST | 14.34 | down | 0.00008 |
| rc_AA179387_at | AA179387 | 379 | DKFZP434N126 protein | 3.7 | down | 0.01588 |
| rc_AA180356_at | AA180356 | 382 | EST | 3.16 | down | 0.00917 |
| rc_AA182030_at | AA182030 | 387 | EST | 8.32 | down | 0.00018 |
| rc_AA182568_at | AA182568 | 388 | STAT induced STAT Inhibitor-2 | 10.92 | down | 0.00099 |
| AA188921_at | AA188921 | 393 | similar to Caenorhabditis elegans protein | 4.05 | down | 0.004 |
| rc_AA190816_at | AA190816 | 395 | EST | 4.1 | down | 0.00037 |
| rc_AA191014_at | AA191014 | 396 | EST | 5.07 | down | 0.01455 |
| ı | | | protein phosphatase 2 (formerly 2A), regulatory | | | |
| rc_AA191310_s_at | AA191310 | 397 | subunit A (PR 65), beta isoform | 7.28 | down | 0 |
| rc_AA191488_s_at | AA191488 | 398 | solute carrier family 31 (copper transporters), | 3.19 | down | 0.00013 |
| rc_AA191647_at | AA191647 | 336 | ceruloplasmin (ferroxidase) | 4.05 | down | 0.00029 |
| rc_AA193204_at | AA193204 | 402 | Arg/Abl-interacting protein ArgBP2 | 8.98 | down | 0.00861 |
| rc_AA193223_at | AA193223 | 403 | EST | 4.2 | down | 0.02416 |
| AA193297_at | AA193297 | 404 | EST | 4.37 | down | 0.04676 |
| rc_AA194075_f_at | AA194075 | 406 | nuclear receptor coactivator 4 | 4.69 | down | 0.00862 |
| AA194146_at | AA194146 | 407 | EST | 3.53 | down | 0.00352 |
| rc_AA194833_at | AA194833 | 411 | claudin 1 | 11.45 | down | 0.00034 |
| rc_AA194997_s_at | AA194997 | 412 | EST . | 16.12 | down | 0.00103 |
| rc_AA195656_at | AA195656 | 418 | KIAA0977 protein | 15.29 | down | 0.00817 |
| rc_AA195657_at | AA195657 | 419 | EST | 6.44 | down | 0.00016 |
| rc_AA196287_at | AA196287 | 420 | EST | 15.07 | down | 0.00001 |
| | | | intercellular adhesion molecule 1 (CD54), human | | | |
| rc_AA197311_s_at | AA197311 | 422 | rhinovirus receptor | 6.07 | down | 0.00053 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| | | | FERM, RhoGEF (ARHGEF) and pleckstrin | | | |
| AA203222_at | AA203222 | 424 | domain protein 1 (chondrocyte-derived) | 4.21 | down | 0.00745 |
| rc_AA205724_at | AA205724 | 426 | EST | 6.73 | down | 0.00027 |
| rc_AA207123_at | AA207123 | 430 | immunoglobulin superfamily, member 3 | 3.16 | down | 0.00328 |
| rc_AA210850_at | AA210850 | 431 | EST | 4.97 | down | 0.00735 |
| rc_AA211388_at | AA211388 | 433 | EST | 4.56 | down | 0.02703 |
| rc_AA214542_at | AA214542 | 438 | EST | 4.43 | down | 0.00601 |
| rc_AA215585 s_at | AA215585 | 442 | nudix (nucleoside diphosphate linked moiety X)- | က | down | 0.03027 |
| AA215919_at | AA215919 | 443 | F-box protein 7 | 6.62 | down | 0.00921 |
| rc_AA218727_at | AA218727 | 445 | EST | 3.47 | down | 0.00125 |
| rc_AA219039_at | AA219039 | 446 | EST | 5.76 | down | 0.00053 |
| rc_AA219304_s_at | AA219304 | 447 | alpha-2-macroglobulin | 21.97 | down | 0.00011 |
| rc_AA219653_at | AA219653 | 448 | EST | 4.08 | down | 0.00607 |
| rc_AA223335_s_at | AA223335 | 449 | propionyl Coenzyme A carboxylase, beta | 5.49 | down | 0.02761 |
| rc_AA223902_at | AA223902 | 450 | EST | 9.91 | down | 0.00003 |
| rc_AA227452_at | AA227452 | 455 | EST | 4.7 | down | 0.02345 |
| rc_AA227480_s_at | AA227480 | 456 | pim-2 oncogene | 3.31 | down | 0.02413 |
| rc_AA227901_at | AA227901 | 459 | SEC24 (S. cerevisiae) related gene family, | 3.18 | down | 0.00397 |
| rc_AA228119_at | AA228119 | 462 | pre-B-cell colony-enhancing factor | 4.77 | down | 0.00031 |
| rc_AA232114_s_at | AA232114 | 463 | epoxide hydrolase 2, cytoplasmic | 24.34 | down | 0.00007 |
| rc_AA233152_at | AA233152 | 467 | EST | 12.95 | down | 0 |
| rc_AA233347_at | AA233347 | 470 | zinc finger protein 216 | 5.9 | down | 0.0041 |
| rc_AA233369_at | AA233369 | 471 | histidine ammonia-fyase | 90.6 | down | 0.0008 |
| rc_AA233763_at | AA233763 | 472 | EST | 4.61 | down | 0.00004 |
| rc_AA233837_at | AA233837 | 474 | EST | 4.79 | down | 0.0034 |
| rc_AA234095_at | AA234095 | 478 | EST | 8.08 | down | 0.00394 |
| rc_AA234527_s_at | AA234527 | 483 | nuclear receptor subfamily 3, group C, member 1 | 6.19 | down | 0.00864 |
| rc_AA234561_at | AA234561 | 485 | EST | 3.88 | down | 0.02058 |
| AA234634_f_at | AA234634 | 486 | CCAAT/enhancer binding protein (C/EBP), delta | 7.48 | down | 0.03318 |
| AA234817_at | AA234817 | 490 | EST | 6.22 | down | 0.00099 |
| rc_AA234831_at | AA234831 | 491 | EST | 3.42 | down | 0.00206 |
| | | | | | | |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|------------|--------|--|-------------|-----------|---------|
| rc AA234916 at | AA234916 | 492 | EST | 3.2 | down | 0.00799 |
| rc_AA235233_at | AA235233 | 493 | EST | 6.59 | down | 0.00755 |
| rc_AA235288_at | AA235288 | 494 | PTPL1-associated RhoGAP 1 | 3.7 | down | 0.00643 |
| rc_AA235310_at | AA235310 | 496 | EST | 37.86 | down | 0.00091 |
| rc_AA235507_at | AA235507 | 498 | golgi autoantigen, golgin subfamily a, 5 | 3.28 | down | 0.00249 |
| rc AA235618 f at | AA235618 | 499 | EST | 3.43 | down | 0.02127 |
| rc_AA235765 s at | AA235765 | 501 | KIAA0214 gene product | 3.59 | down | 0.01148 |
| rc_AA235811_at | AA235811 | 502 | EST | 3.64 | down | 0.01272 |
| rc AA235873 s at | AA235873 | 505 | H factor (complement)-like 1,H factor 1 | 9.98 | down | 0.01667 |
| rc_AA236230_at | · AA236230 | 508 | EST | 5.28 | down | 0.01517 |
| | AA236365 | 209 | 3-phosphoglycerate dehydrogenase | 10.23 | down | 0.00562 |
| rc AA236401 at | AA236401 | 510 | EST | 16.71 | down | 0.00088 |
| rc AA236455 r at | AA236455 | 512 | EST | 15.71 | down | 0.00286 |
| rc_AA236455_s_at | AA236455 | 512 | EST | 11.35 | down | 0.02859 |
| rc_AA236796 s at | AA236796 | 517 | follistatin | 8.74 | down | 0.00862 |
| rc AA236942 at | AA236942 | 519 | EST | 3.18 | down | 0 |
| rc_AA236982_at | AA236982 | 520 | sterol carrier protein 2 | 5.56 | down | 0.01542 |
| rc_AA242766_at | AA242766 | 523 | EST | 3.58 | down | 0.0151 |
| rc_AA243495_at | AA243495 | 528 | lectin, mannose-binding, 1 | 4.23 | down | 0.00179 |
| rc_AA243582_at | AA243582 | 529 | hemoglobin, gamma A | 7.15 | down | 0.0021 |
| rc_AA243595 s at | AA243595 | 530 | EST | 3.11 | down | 0.008 |
| AA247453 at | AA247453 | 533 | EST | 3.09 | down | 0.0015 |
| rc_AA250744_at | AA250744 | 536 | EST | 3.39 | пмор | 0.01137 |
| rc_AA250775_at | AA250775 | 537 | EST | 4.52 | down | 0.01752 |
| rc_AA251114_at | AA251114 | 539 | prostate cancer overexpressed gene 1 | 9.9 | down | 0.00039 |
| rc_AA251837_at | AA251837 | 547 | EST | 3.87 | down | 0.00782 |
| | | | quinolinate phosphoribosyltransferase (nicotinate- | | | |
| rc_AA252289_at | AA252289 | 552 | nucleotide pyrophosphorylase (carboxylating)) | 5.66 | down | 0.01389 |
| rc_AA252365_at | AA252365 | 554 | EST | 3.9 | down | 0.01796 |
| rc_AA253043_at | AA253043 | 559 | DKFZP586I1419 protein | 3.89 | down | 0.00145 |
| AA253129_at | AA253129 | 260 | F-box protein FBL11 | 6.47 | down | 0.00001 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|----------------------------|-------------|-----------|---------|
| rc_AA253216_at | AA253216 | 561 | EST | 28.18 | | 0.00141 |
| rc_AA253369_s_at | AA253369 | 563 | EST | 15.59 | down | 0.00091 |
| rc_AA253455_s_at | AA253455 | 565 | EST | 3.05 | down | 0.00533 |
| rc AA253459 at | AA253459 | 566 | EST | 4.51 | down | 0.00419 |
| rc_AA255546_at | AA255546 | 569 | EST | 4 | down | 0.00301 |
| rc_AA255624_at | AA255624 | 571 | EST | 4.06 | down | 69000.0 |
| rc_AA255878_at | AA255878 | 572 | KIAA0767 protein | 3.96 | down | 0.00592 |
| rc_AA255903_at | AA255903 | 573 | CD39-like 4 | 2.67 | down | 0.01687 |
| rc_AA256171_at | AA256171 | 575 | EST | 7.34 | down | 0.04562 |
| rc_AA256341_at | AA256341 | 578 | EST | 7.37 | down | 0.00091 |
| rc_AA256367_s_at | AA256367 | 579 | paraoxonase 3 | 70.33 | down | 0.00192 |
| rc_AA256666_at | AA256666 | 583 | EST | 4.63 | down | 0.0018 |
| rc_AA257057_s_at | AA257057 | 586 | EST | 8.11 | down | 0.00379 |
| rc_AA258308_at · | AA258308 | 290 | EST | 5.4 | down | 0.00023 |
| rc_AA258323_at | AA258323 | 591 | EST | 4.31 | down | 0.00046 |
| rc_AA258350_at | AA258350 | 592 | EST | 5.08 | down | 0.00035 |
| rc_AA258353_at | AA258353 | 593 | EST | 5.28 | down | 0.00193 |
| rc_AA258567_at | AA258567 | 297 | EST | 6.92 | down | 96000.0 |
| rc_AA258613_at | AA258613 | 598 | EST | 4.31 | down | 0.0344 |
| rc_AA258813_at | AA258813 | 900 | EST | 4.63 | down | 0.02395 |
| rc_AA259064_at | AA259064 | 602 | EST | 13.15 | down | 0.00001 |
| rc_AA261954_at | AA261954 | 604 | EST | 7.69 | down | 0.00334 |
| rc_AA262033_s_at | AA262033 | 909 | EST | 4.41 | down | 0.00054 |
| rc_AA262349_at | AA262349 | . 607 | EST | 3.78 | down | 0.00043 |
| rc_AA262766_at | AA262766 | 609 | EST | 99.5 | down | 0.03832 |
| rc_AA279112_at | AA279112 | 622 | EST | 3.42 | down | 0.01444 |
| rc_AA279533_at | AA279533 | 627 | EST | 5.01 | down | 0.04448 |
| AA279550_at | AA279550 | 628 | Kruppel-like factor | 4.06 | down | 0.00957 |
| rc_AA279676_s_at | AA279676 | 630 | deoxyribonuclease I-like 3 | 23.35 | down | 0.00001 |
| rc_AA279802_at | AA279802 | 631 | EST | 3.65 | down | 0.03366 |
| rc_AA279937_at | AA279937 | 634 | EST | 3.38 | down | 0.02719 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affv ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| rc AA280130 at | AA280130 | 636 | EST | 4.12 | down | 0.00114 |
| 1 | | | spleen focus forming virus (SFFV) proviral | | | |
| rc AA280413 s at | AA280413 | 638 | integration oncogene spi1 | 4.46 | down | 0.02062 |
| rc AA280791 at | AA280791 | 640 | eukaryotic translation initiation factor 5 | 3.11 | down | 0.03339 |
| rc_AA281440_at | AA281440 | 644 | EST | 6.43 | down | 0.01246 |
| rc_AA281545_at | AA281545 | 645 | EST | 3.64 | down | 0.00002 |
| rc_AA281591_at | AA281591 | 646 | EST | 3.23 | down | 0.00895 |
| | AA281677 | 648 | DKFZP564M2423 protein | 3.95 | down | 0.03606 |
| rc AA281770 at | AA281770 | 649 | seven in absentia (Drosophila) homolog 1 | 3.96 | down | 0.00094 |
| rc_AA281796_at | AA281796 | . 650 | mannose-P-dolichol utilitzation defect 1 | 3.3 | down | 0.04108 |
| ı | | | core-binding factor, runt domain, alpha subunit 2; | • | | |
| rc AA281930 at | AA281930 | 651 | translocated to, 3 | 3.27 | down | 0.02329 |
| rc_AA282061_at | AA282061 | 652 | KIAA0962 protein | 8.95 | down | 0.01033 |
| rc_AA282089_at | AA282089 | 653 | EST | 4.93 | down | 0.00108 |
| rc_AA282179_at | AA282179 | 655 | EST . | 3.09 | down | 0.01693 |
| rc_AA282238_at | AA282238 | 656 | EST | 3.47 | down | 0.00677 |
| rc_AA282516_at | AA282516 | 099 | 7-dehydrocholesterol reductase | 7.67 | down | 0.0008 |
| rc_AA282886_at | AA282886 | 663 | EST | 3.57 | down | 0.00049 |
| rc_AA282971_at | AA282971 | 665 | EST | 4.37 | down | 0.03822 |
| rc_AA283758_at | AA283758 | 670 | EST | 3.67 | down | 0.04293 |
| AA284558 at | AA284558 | 674 | Nck, Ash and phospholipase C binding protein | 3.09 | down | 0.00027 |
| rc aa284721_s at | AA284721 | 677 | EST | 3.34 | down | 0.03296 |
| rc AA284795_at | AA284795 | 678 | phosphatidylethanolamine N-methyltransferase | 10.03 | down | 0.00019 |
| rc_AA285053_at | AA285053 | 681 | EST | 6.95 | down | 0.00125 |
| rc_AA287122_at | AA287122 | 989 | EST | 3.66 | down | 0.00161 |
| rc AA287550 f at | AA287550 | 689 | DKFZP434C171 protein | 3.53 | down | 0.00217 |
| rc_AA287566_at | AA287566 | 069 | KIAA0187 gene product | 9.07 | down | 0.00013 |
| rc_AA291323_at | AA291323 | 669 | BCL2-interacting killer (apoptosis-inducing) | 9.15 | down | 0.00514 |
| rc AA291749 s at | AA291749 | 703 | estrogen receptor 1 | 4.78 | down | 0.00059 |
| rc_aa292086_s_at | AA292086 | 705 | EST | 2 | down | 0.00161 |
| AA292158_s_at | AA292158 | 200 | EST | 21.79 | down | 0.00031 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| rc_AA292328_at | AA292328 | 707 | activating transcription factor 5 | 17.51 | down | 0.00689 |
| rc_AA292711_at | AA292711 | 711 | EST | თ | down | 0.01053 |
| rc AA292773 s at | AA292773 | 713 | collagen, type XVIII, alpha 1 | 7.44 | down | 0.00158 |
| rc_AA293327_at | AA293327 | 716 | isocitrate dehydrogenase 1 (NADP+), soluble | 7.04 | down | 0.04377 |
| rc_AA293485_at | AA293485 | 718 | EST | 3.36 | down | 0.02799 |
| AA298180_at | AA298180 | 726 | EST | 3.11 | down | 0.00747 |
| rc_AA299632_at | AA299632 | 728 | EST | 4.23 | down | 0.00371 |
| rc_AA312946_s_at | AA312946 | 731 | EST | 9.21 | down | 0.00106 |
| AA314457_at | AA314457 | 733 | synaptonemal complex protein 3 | 4.86 | down | 0.0013 |
| rc_AA338512_at | AA338512 | 742 | EST | 3.05 | down | 0.03427 |
| rc_AA342301_at | AA342301 | 746 | EST | 3.89 | down | 0.00038 |
| rc_AA342337_at | AA342337 | 747 | EST | 3.87 | down | 0.0069 |
| rc_AA342446_s_at | AA342446 | 748 | insulin receptor | 6.83 | down | 0.00412 |
| rc_AA342771_at | AA342771 | 749 | EST | 5.33 | down | 0.00331 |
| rc_AA343142_at | AA343142 | 751 | EST | 20.87 | down | 0.00003 |
| rc_AA344866_s_at | AA344866 | 752 | complement component 8, gamma polypeptide | 7.28 | down | 0.00206 |
| rc_AA347674_at | AA347674 | 753 | EST | 10.59 | down | 0.03716 |
| rc_AA347717_at | AA347717 | 754 | EST | 5.25 | down | 0.00207 |
| rc_AA348284_at | AA348284 | 755 | EST | 4.54 | down | 0.00759 |
| rc_AA348466_s_at | AA348466 | 756 | regulator of G-protein signalling 5 | 3.2 | down | 0.00571 |
| rc_AA348485_at | AA348485 | 757 | KIAA0438 gene product | 4.01 | down | 0.04563 |
| i | | | fatty-acid-Coenzyme A ligase, long-chain 1, fatty- | | | • |
| rc_AA348922_s_at | AA348922 | 758 | acid-Coenzyme A ligase, long-chain 2 | 64.27 | down | 0.00002 |
| rc_AA349836_at | AA349836 | 760 | EST | 3.01 | - down | 0.00911 |
| | | | KIAA0382 protein; leukemia-associated rho | | | |
| rc_AA370359_s_at | AA370359 | 767 | guanine nucleotide exchange factor (GEF) | 4.82 | down | 0.01077 |
| AA376875_at | AA376875 | 770 | monoamine oxidase A | 3.8 | down | 0.02746 |
| rc_AA377087_at | AA377087 | 771 | EST | 16.75 | down | 0.00002 |
| rc_AA381125_at | AA381125 | 772 | EST | 15.48 | down | 0 |
| rc_AA382975_f_at | AA382975 | 773 | EST | 3.7 | down | 0.00131 |
| rc_AA393825_at | AA393825 | 776 | EST | 3.62 | down | 0.0065 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| AA393961_at | AA393961 | 777 | EST | 3.77 | down | 0.01029 |
| AA397841_at | AA397841 | 780 | EST | 8.21 | down | 0 |
| rc_AA397914_at | AA397914 | 783 | EST | 3.16 | down | 0.00336 |
| rc_AA398102_at | AA398102 | 786 | KIAĄ0429 gene product | 6.22 | down | 0.00396 |
| rc_AA398124_s_at | AA398124 | 787 | growth factor receptor-bound protein 14 | 7.82 | down | 60000.0 |
| rc_AA398221_at | AA398221 | 790 | calcium/calmodulin-dependent protein kinase | 3.78 | down | 0.00019 |
| rc_AA398257_at | AA398257 | 791 | 7-dehydrocholesterol reductase | 4.43 | down | 0.04169 |
| rc_AA398280_at | AA398280 | 792 | EST | 12.43 | down | 0.00134 |
| rc_AA398386_at | AA398386 | 793 | EST | 5.71 | down | 0.00007 |
| rc_AA398422_i_at | AA398422 | 794 | EST | 3.94 | down | 0.00388 |
| rc_AA398423_at | AA398423 | 795 | EST | 8.26 | down | 0.00063 |
| rc_AA398445_at | AA398445 | 796 | EST | 4.28 | down | 0.01764 |
| rc_AA398892_at | AA398892 | 800 | similar to yeast BET3 (S. cerevislae) | 7.43 | down | 0.00038 |
| rc_AA400030_at | AA400030 | 806 | EST | 3.98 | down | 0.00088 |
| rc_AA400246_at | AA400246 | 810 | mitogen-activated protein kinase-activated | 3.09 | down | 0.00476 |
| rc_AA400251_at | AA400251 | 811 | EST | 4.07 | down | 0.00032 |
| rc_AA400258_at | AA400258 | 812 | EST | 11.89 | down | 0.00478 |
| rc_AA400259_at | AA400259 | 813 | EST | 3.65 | down | 0.00476 |
| rc_AA400471_at | AA400471 | 816 | EST | 5.45 | down | 0.0056 |
| rc_AA400780_at | AA400780 | 818 | EST | 3.5 | down | 0.00107 |
| rc_AA400831_at | AA400831 | 819 | EST | 3.49 | down | 0.00105 |
| rc_AA400834_f_at | AA400834 | 820 | EST | 4.73 | down | 0.01523 |
| rc_AA400864_at | AA400864 | 821 | EST | 7.51 | down | 0.02237 |
| rc_AA400915_at | AA400915 | 823 | EST | 9.84 | down | 0.00351 |
| rc_AA400934_at | AA400934 | 824 | EST | 4.98 | down | 0.02013 |
| rc_AA400979_at | AA400979 | 825 | calcitonin receptor-like receptor activity modifying | 6.65 | down | 0.01051 |
| rc_AA401151_at | AA401151 | 827 | lysozyme (renal amyloidosis) | 3.01 | down | 0.0051 |
| rc_AA401343_at | AA401343 | 828 | EST | 3.11 | down | 0.01929 |
| rc_AA401376_at | AA401376 | 829 | EST | 3.97 | down | 0.00797 |
| rc_AA401562_s_at | AA401562 | 830 | EST | 50.45 | down | 0.00301 |
| AA402006_at | AA402006 | 834 | EST | 4.19 | down | 0.00094 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| AA402095 s_at | AA402095 | 835 | EST | 3.12 | down | 0.01412 |
| rc_AA402224_at | AA402224 | 836 | growth arrest and DNA-damage-inducible, | 14.41 | down | 0.00012 |
| rc_AA402656_at | AA402656 | 841 | EST | 12.05 | down | 0.00001 |
| rc_AA402799_at | AA402799 | 842 | EST | 11.81 | down | 0.00031 |
| AA404252 at | AA404252 | 848 | lectin, mannose-binding, 1 | 16.15 | down | 0.00001 |
| rc_AA404352_at | AA404352 | 820 | EST | 7 | down | 0.00059 |
| rc_AA404500 at | AA404500 | 852 | EST | 4.16 | down | 0.01375 |
| rc_AA405819_at | AA405819 | 865 | KIAA0668 protein | 8.59 | down | 0.02034 |
| rc_AA405832_at | AA405832 | 998 | EST | 12.24 | down | 0.00441 |
| rc_AA405907_at | AA405907 | 867 | EST | 3.12 | down | 0 |
| rc_aa406125_s_at | AA406125 | 898 | EST | 4.95 | down | 0.01027 |
| rc_AA406126_at | AA406126 | 869 | ÉST | 8.43 | down | 0.00569 |
| rc_AA406231_s_at | AA406231 | 873 | KIAA0381 protein | 4.46 | down | 0.04049 |
| AA406435_at | AA406435 | 877 | EST | 3.24 | down | 0.00941 |
| rc_AA410181_at | AA410181 | 881 | EST | ဇ | down | 0.00268 |
| rc_AA410255_at | AA410255 | 882 | EST | 7.56 | down | 0.00043 |
| rc_AA410507_at | AA410507 | 884 | EST | 3.73 | down | 0.01703 |
| rc_AA410523_at | AA410523 | 886 | EST | 6.37 | down | 0.03506 |
| rc_AA411764_at | AA411764 | 891 | similar to APOBEC1 | 4 | down | 0.01491 |
| rc_AA412034_at | AA412034 | 894 | EST | 3.09 | down | 0.02309 |
| rc_AA412063_at | AA412063 | 895 | EST | 8.26 | down | 0.00001 |
| rc_AA412184_at | AA412184 | 888 | EST | 3.08 | down | 0.00012 |
| rc_AA412481_s_at | AA412481 | 905 | EST | 8.07 | down | 0.00014 |
| rc_AA416723_at | AA416723 | 906 | EST | 3.57 | down. | 0.01042 |
| rc_AA416740_at | AA416740 | 206 | EST | 3.08 | down | 0.01592 |
| rc_AA416873_at | AA416873 | 908 | EST | 7.82 | down | 0.00005 |
| | | | sphingomyelin phosphodiesterase 1, acid | | | |
| rc_AA416890_s_at | AA416890 | 606 | lysosomal (acid sphingomyelinase) | 9.9 | down | 0.00112 |
| | | | 5-methyltetrahydrofolate-homocysteine | | | |
| rc_AA416936_at | AA416936 | 910 | methyltransferase reductase | 4.98 | down | 0.00632 |
| rc_AA417046_at | AA417046 | 915 | fatty-acid-Coenzyme A ligase, very long-chain 1 | 44 | down | 0 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| rc AA417078 at | AA417078 | 916 | EST | 4.1 | down | 0.00414 |
| rc_AA417373_at | AA417373 | 917 | EST | 4.8 | down | 0.01342 |
| rc_AA417375_at | AA417375 | 918 | EST | 4.21 | down | 0.00231 |
| AA418098 at | AA418098 | 920 | cAMP responsive element binding protein-like 2 | 3.18 | down | 0.03824 |
| I | | | cytochrome P450, subfamily I (aromatic | | | |
| rc AA418907 s at | AA418907 | 922 | compound-inducible), polypeptide 1 | 4.05 | down | 0.04276 |
| | AA419507 | 924 | EST | 5.58 | down | 0.00578 |
| rc AA419608 at | AA419608 | 925 | EST | 9.19 | down | 0.00005 |
| rc_AA419622_at | AA419622 | 926 | EST | 4.62 | down | 0.00386 |
| rc_AA421049_at | AA421049 | 927 | activating transcription factor 5 | 44.41 | down | 0.00179 |
| rc_AA421052_at | AA421052 | 929 | branched chain alpha-ketoacid dehydrogenase | 3.52 | down | 0.00869 |
| rc AA421244 s at | AA421244 | 932 | SH3-domain binding protein 5 (BTK-associated) | 4.32 | down | 0.007 |
| rc AA421561 at | AA421561 | 933 | insulin-like growth factor 2 (somatomedin A) | 96.6 | down | 0.00007 |
| AA424307 at | AA424307 | 944 | EST | 5.73 | down | 0.0074 |
| rc_AA424672_s at | AA424672 | 946 | dermatopontin | 4.69 | down | 0.00843 |
| | AA424798 | 947 | EST | 17.45 | down | 0.00352 |
| rc_AA424813_at | AA424813 | 948 | EST | 5.77 | down | 0.00503 |
| rc_AA425294_at | AA425294 | 952 | EST | 10.61 | down | 0.00083 |
| rc_AA425309_at | AA425309 | 953 | nuclear factor I/B | 4.9 | down | 0.00466 |
| rc_AA425782_at | AA425782 | 926 | KIAA0874 protein | 5.52 | down | 0.03433 |
| rc_AA425836_at | AA425836 | 957 | EST | 4.55 | down | 0.00035 |
| AA426156_at | AA426156 | 929 | EST | 3.67 | down | 0.00153 |
| AA426168 at | AA426168 | 096 | KIAA0805 protein | 3.73 | down | 0.01477 |
| AA426304 s at | AA426304 | 962 | EST | 6.61 | down | 0.01092 |
| rc_AA426330_at | AA426330 | 963 | N-acylsphingosine amidohydrolase (acid | 4.24 | down | 0.00668 |
| rc_AA426468_at | AA426468 | 996 | EST | 3.38 | down | 0.0099 |
| rc_AA426609_at | AA426609 | 896 | EST | 6.28 | down | 0.01233 |
| rc AA427778 at | AA427778 | 978 | EST | 3.57 | down | 0.00368 |
| AA427783 at | AA427783 | 979 | EST | 4.37 | down | 0.0004 |
| rc_AA427819_at | AA427819 | 980 | midline 2 | 3.44 | down | 0.00063 |
| AA428006_at | AA428006 | 984 | DKFZP564B167 protein | 3.71 | down | 0.02325 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| at AA428150 985 at AA428325 988 last AA428567 989 last AA428607 990 last AA428900 992 last AA428900 992 last AA429904 1005 last AA429004 1005 last AA430028 1008 last AA430028 1008 last AA430047 1011 last AA431462 1022 last AA431480 1023 last AA43591 1038 last AA43591 1038 last AA43591 1038 last AA435824 1045 last AA435824 1045 last AA435824 1053 last AA435885 1049 last AA43680 1053 last AA43680 1055 last AA43680 last | Genbank Seq ID Known Gene Name Fo | Fold Change | Direction | Pvalue |
|--|--|-------------|-----------|---------|
| AA428325 988 1 AA42867 989 1 AA428607 990 1 AA428900 992 1 AA428900 992 1 AA429904 1005 1 AA429904 1005 1 AA430026 1007 1 AA430026 1007 1 AA430044 1010 1 AA430044 1010 1 AA430046 1002 1 AA43066 1016 1 AA43066 1016 1 AA431773 1026 1 AA431773 1026 1 AA435591 1038 1 AA435591 1038 1 AA435891 1038 1 AA435891 1038 1 AA435894 1039 1 AA435894 1039 1 AA435894 1053 1 AA43689 1053 1 | 982 | 5.24 | down | 0.00167 |
| AA42867 989 1 AA428607 990 1 AA428607 990 1 AA428008 992 1 AA429038 991 1 AA429038 992 1 AA429038 995 1 AA429038 995 1 AA42904 1005 1 AA430011 1006 1 AA430026 1007 1 AA430047 1011 1 AA430047 1011 1 AA430048 1013 1 AA431480 1023 1020 1 AA431773 1026 1038 1 AA435591 1038 1045 11 AA435891 1038 1045 11 AA435885 1049 11053 1 | 886 | 8.36 | down | 0.00002 |
| AA428607 990 1 AA428863 991 1 AA428900 992 1 AA429038 995 1 AA429038 995 1 AA429038 995 1 AA42904 1005 1 AA430011 1006 1 AA430026 1007 1 AA430044 1010 1 AA430047 1011 1 AA430108 1013 1 AA431773 1020 1023 1 AA431773 1026 1038 1 AA435591 1038 1 AA435591 1038 1 AA435885 1045 11 AA435885 1049 11053 1 | 686 | 3.99 | down | 0.00788 |
| AA428863 991 1 AA428900 992 1 AA429038 995 1 AA429038 995 1 AA42904 1005 1 AA430011 1006 1 AA430026 1007 1 AA43004 1010 1 AA43004 1010 1 AA43004 1010 1 AA430066 1016 1 AA431773 1020 1 AA431773 1026 1 AA435591 1038 1 AA435591 1038 1 AA43588 1045 1 AA43588 1045 1 AA43588 1049 11053 1 | | 4.21 | down | 0.04305 |
| AA428900 992 But AA429038 995 But AA429038 995 But AA429904 1005 But AA430026 1007 But AA430028 1008 But AA43004 1010 But AA43004 1010 But AA430066 1016 But AA431773 1026 But AA435591 1038 But AA435777 1047 But AA435824 1048 But AA435824 1049 But AA435824 1055 But AA43680 1055 | 991 | 3.5 | down | 0.01726 |
| at AA429038 995 at AA429478 998 at AA429904 1005 at AA430026 1007 at AA430028 1008 at AA430028 1008 at AA430028 1022 at AA431773 1020 at AA435591 1033 at AA435777 1047 at AA435824 1048 at AA435895 1049 at AA436489 1053 at AA436489 1053 at AA436489 1053 at AA436489 1053 at AA436489 1055 at AA436560 1056 | 992 | 7.01 | down | 0.00037 |
| at AA429478 998 1 AA429904 1005 1 AA430011 1006 1 AA430028 1008 1007 1 AA430044 1010 1 AA430047 1011 102 AA430108 1013 102 AA431773 1020 1023 1044 AA435591 1033 1045 AA435777 1047 AA435824 1048 AA435824 1048 AA435824 1048 AA435660 1055 | 995 | 3.29 | down | 0.00927 |
| at AA429904 1005 AA430011 1006 AA430026 1007 AA430028 1008 AA430044 1010 AA430044 1010 AA430047 1011 AA430047 1011 AA430047 1011 AA430047 1011 AA430047 1011 AA430108 1013 AA431773 1026 AA431773 1026 AA431773 1026 AA435591 1038 AA435777 1047 AA435824 1048 AA435824 1048 AA435824 1048 AA435824 1053 AA435660 1055 | 866 | 3.41 | down | 0.02599 |
| AA430011 1006 AA430026 1007 AA430028 1008 AA430044 1010 AA430047 1010 AA430047 1011 AA430047 1011 AA430047 1011 AA431337 1020 AA431462 1022 AA431462 1022 AA431473 1026 AA431480 1023 AA431773 1026 AA432168 1033 AA435591 1038 AA435591 1038 AA435591 1045 AA435824 1048 AA435824 1048 AA435824 1048 AA435885 1049 | 1005 | 7.26 | down | 0.00524 |
| at AA430026 1007 at AA430028 1008 at AA430044 1010 at AA430047 1011 at AA430108 1013 at AA431337 1020 at AA431462 1022 at AA431480 1023 at AA432168 1031 at AA43591 1038 at AA435777 1047 at AA435824 1048 at AA436489 1053 at AA436489 1053 | 1006 | 8.35 | down | 0.00729 |
| at AA430028 1008 at AA430044 1010 at AA430047 1011 at AA430047 1011 at AA431337 1020 at AA431462 1022 at AA431480 1023 at AA43591 1038 at AA43591 1045 at AA435824 1049 at AA435824 1048 at AA436860 1055 at AA436800 105 | 1007 | 3.31 | down | 0.00786 |
| at AA430044 1010 AA430047 1011 AA430048 1013 AA430108 1013 AA431337 1020 AA431480 1023 AA431480 1023 AA431480 1023 AA431773 1026 AA431773 1026 AA435591 1038 AA435591 1045 AA435885 1049 AA435885 1049 AA435885 1049 AA435885 1049 AA435885 1049 AA435885 1055 AA436860 1055 AA436560 1055 AA436560 1055 | 1008 | 9.14 | down | 0.00246 |
| at AA430047 1011 at AA430108 1013 at AA431337 1020 at AA431462 1022 at AA431473 1026 at AA431773 1026 at AA432168 1033 at AA435591 1038 at AA435591 1045 at AA435824 1045 at AA435824 1045 at AA436860 1053 | 1010 | 7.78 | down | 0.00124 |
| at AA430108 1013 at AA430666 1016 at AA431337 1020 at AA431480 1023 at AA431773 1026 at AA432168 1033 at AA435591 1038 at AA435777 1047 at AA435824 1048 at AA436860 1055 at AA436860 1055 at AA436560 1055 at AA45660 1055 at AA45600 1055 at AA45600 1055 at AA45600 1055 at AA56600 1 | 1011 | 3.44 | down | 0:0016 |
| at AA430666 1016 1016 at AA431337 1020 at AA431462 1022 at AA431773 1026 at AA432168 1031 at AA435591 1038 at AA435757 1047 at AA435824 1049 at AA436489 1053 at AA436560 1055 at | 1013 | 3.8 | down | 0.04484 |
| at AA431337 1020 at AA431462 1022 at AA431480 1023 at AA431773 1026 at AA432168 1031 at AA43591 1038 at AA43577 1047 at AA435985 1049 at AA436985 1055 at AA436560 1055 | 1016 | 5.12 | down | 0.00377 |
| at AA431462 1022 AA431480 1023 at AA431773 1026 at AA432168 1031 at AA435591 1038 at AA435753 1045 at AA435824 1048 at AA435824 1048 at AA436489 1053 at AA436560 1055 | 1020 | 6.26 | down | 0.00053 |
| at AA431480 1023 at AA431773 1026 at AA432168 1031 at AA435591 1038 at AA435753 1045 at AA435777 1047 at AA435824 1048 at AA436489 1053 at AA436560 1055 | 1022 | 4.45 | down | 0.00956 |
| at AA431773 1026 at AA432168 1031 at AA433946 1033 at AA435591 1038 at AA435777 1047 at AA435824 1048 at AA436489 1053 at AA436560 1055 | 1023 | 4.3 | down | 0.00876 |
| AA432168 1031 AA433946 1033 AA435753 1045 AA435777 1047 AA435824 1048 AA435985 1049 AA436489 1053 AA436560 1055 | 1026 | 7.61 | down | 0.00063 |
| AA43546 1033 AA435591 1038 AA43577 1045 AA435824 1048 AA435985 1049 AA436489 1053 AA43650 1055 | 1031 | 4.71 | down | 0.01377 |
| AA435753 1045 AA435777 1047 AA435824 1048 AA435985 1049 AA436489 1053 AA436560 1055 | 1033 | 43.74 | down | 0.00005 |
| AA435753 1045 AA435777 1047 AA435985 1049 AA436489 1053 AA436560 1055 | | 3.5 | down | 0.0001 |
| AA435777 1047 AA435824 1048 AA435985 1049 AA436489 1053 AA436560 1055 | 1045 | 4.71 | down | 0.00078 |
| AA435824 1048 AA435985 1049 AA436489 1053 AA436560 1055 | solute carrier family 25 (mitochondrial carrier; | | | |
| AA435824 1048 AA435985 1049 AA436489 1053 AA436560 1055 | • | 7.48 | down | 0.00613 |
| at AA435985 1049 at AA436489 1053 at AA436560 1055 | 1048 | 3.93 | down | 0.02764 |
| at AA436489 1053 at AA436560 1055 | 1049 | 17.7 | down | 0 |
| _at AA436560 1055 | 1053 | 7.34 | down | 0.001 |
| | 1055 | 11.41 | down | 0.00756 |
| | 3690 1057 EST | 4.58 | down | 0.00948 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| rc AA436880 at | AA436880 | 1058 | EST | 3.22 | down | 0.00699 |
| AA436926 at | AA436926 | 1059 | EST | 5.5 | down | 0.00984 |
| rc AA437235 s at | AA437235 | 1060 | EST | 7.15 | down | 0.01455 |
| AA437265_s | AA437265 | 1061 | EST | 4.39 | down | 0.00826 |
| rc AA437295 at | AA437295 | 1062 | ribosomal protein L7a | 4.35 | down | 0.00347 |
| rc_AA441791_at | AA441791 | 1065 | EST | 3.58 | down | 0.00357 |
| AA442334 at | AA442334 | 1069 | EST | 7.15 | down | 0.00018 |
| AA442342 at | AA442342 | 1070 | EST | 5.62 | down | 0.00052 |
| rc_AA443272_at | AA443272 | 1074 | EST | 7.68 | down | 0.00869 |
| rc_AA443658_at | AA443658 | 1079 | transmembrane 7 superfamily member 2 | 90.6 | down | 0.00048 |
| rc_AA443756_at | AA443756 | 1080 | EST | 5.05 | down | 0.00341 |
| rc_AA443822_at | AA443822 | 1082 | EST | 5.46 | down | 0.02538 |
| rc_AA443934_at | AA443934 | 1083 | GTP-binding protein Rho7 | 3.09 | down | 0.00214 |
| rc AA443936 s at | AA443936 | 1084 | EST | 22.96 | down | 0.00627 |
| rc AA443993 at | AA443993 | 1086 | EST | 3.21 | down | 0.02948 |
| rc_AA446342_at | AA446342 | 1088 | seven in absentia (Drosophila) homolog 1 | 4.84 | down | 0.00015 |
| rc_AA446587_at | AA446587 | 1091 | EST | 5.8 | down | 0.00012 |
| rc_AA446651_at | AA446651 | 1093 | EST | 3.14 | down | 0.01902 |
| rc_AA446666_at | AA446666 | 1094 | EST | 4.03 | down | 0.02369 |
| 1 | | | UDP-N-acteylglucosamine pyrophosphorylase 1; | | | |
| rc_AA447549_at | AA447549 | 1101 | Sperm associated antigen 2 | 6.37 | down | 0.02815 |
| rc_AA447617_at | AA447617 | 1103 | EST | 3.26 | down | 0.04687 |
| rc_AA447740_at | AA447740 | 1106 | EST | 3.22 | down | 0.02518 |
| rc_AA447971_at | AA447971 | 1110 | EST | 8.08 | down | 0.00035 |
| rc_AA447977_s_at | AA447977 | 1111 | EST | 3.84 | down | 0.00045 |
| rc AA448002 at | AA448002 | 1113 | putative type II membrane protein | 10.05 | down | 0 |
| rc_AA448282_at | AA448282 | 1115 | EST | 3.87 | down | 0.00217 |
| | , | | FXYD domain-containing ion transport regulator 1 | | | |
| rc_AA448300_at | AA448300 | 1116 | (phospholemman) | 24.97 | down | 0.00001 |
| rc_AA449267_at | AA449267 | 1120 | EST | 16.44 | down | 0.00926 |
| | AA449297 | 1121 | EST | 3.78 | down | 0.00039 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| | | 2 | MISWII SCHO HAIRE | I Old Ollalige | חופפוום | 2 |
|------------------|------------|------|--|----------------|---------|---------|
| rc_AA449306_at | AA449306 | 1122 | EST | 5.01 | down | 0.0006 |
| rc_AA449327_at | AA449327 | 1123 | EST | 4.77 | down | 0.01248 |
| | AA449448 | 1125 | EST | 3.97 | down | 0.00103 |
| rc AA450114 at | AA450114 | 1131 | EST | 4.96 | down | 0.01238 |
| rc_AA450127_at | AA450127 | 1132 | growth arrest and DNA-damage-inducible, beta | 7.98 | down | 0.00078 |
| rc_AA450281_at | AA450281 | 1134 | EST | 5.55 | down | 0.00004 |
| rc_AA451836_at | AA451836 | 1137 | EST | 4.9 | down | 0.01412 |
| rc AA451911 at | AA451911 | 1139 | EST | 3.44 | down. | 0.00221 |
| rc_AA452158_at | AA452158 | 1141 | ras homolog gene family, member B | 28.96 | down | 0.00064 |
| AA452454_at | AA452454 | 1144 | EST | 4.45 | down | 0.00179 |
| rc AA452549 at | AA452549 | 1146 | platelet-derived growth factor receptor, alpha | 3.3 | down | 0.04155 |
| rc_AA452559_s_at | AA452559 | 1147 | EST | 4.35 | down | 0.00804 |
| rc_aa452598_s_at | AA452598 | 1148 | genethonin 1 | 5.49 | down | 0.00163 |
| rc AA452855 at | AA452855 | 1150 | lectin, mannose-binding, 1 | 9.88 | down | 0.00428 |
| rc_AA452860_at | AA452860 | 1151 | EST | 3.99 | down | 0.00831 |
| rc_AA452915_at | AA452915 | 1152 | EST | 3.13 | down | 0.00561 |
| rc_AA453770_s_at | AA453770 | 1157 | EST | 6.04 | down | 0.00524 |
| AA453917 at | AA453917 | 1159 | EST | 3.3 | down | 0.01896 |
| rc AA453988 at | AA453988 | 1160 | methionine adenosyltransferase I, alpha | 54.29 | down | 0.00381 |
| rc_AA454086_f_at | AA454086 | 1161 | UDP-glucose dehydrogenase | 4.29 | down | 0.00981 |
| rc AA454159_at | AA454159 | 1162 | EST | 10.81 | down | 0.00132 |
| rc_AA454170_at | AA454170 | 1163 | EST | 3.11 | down | 0.03 |
| rc AA454177 i at | AA454177 | 1164 | EST | 10.3 | down | 0.0008 |
| rc AA454184 at | AA454184 | 1165 | EST | 3.96 | down | 0.04605 |
| rc_AA454733_s_at | AA454733 | 1169 | EST | 5.61 | down | 0.01182 |
| rc_AA455097 i at | AA455097 | 1172 | EST | 6.03 | down | 0.00419 |
| rc_AA455367_at | AA455367 | 1176 | DKFZP586F1018 protein | 3.73 | down | 0.00202 |
| AA455403_at | AA455403 | 1177 | EST | 15.46 | down | 0.01547 |
| rc_AA455865_at | AA455865 | 1180 | phosphatidylinositol glycan, class B | 5.41 | down | 0.00004 |
| rc AA455896 s_at | AA455896 | 1181 | glypican 1 | 3.46 | down | 0.00887 |
| A A AEEOED OF | A A AEEOGO | 1100 | TOT | • | 40,000 | 20000 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| | AA455987 AA456988 AA456055 AA456147 AA456289 AA4562811 | 1183 | EST butyrobetaine (gamma), 2-oxoglutarate | 5.36 | down | 0.00029 |
|--|---|------|---|-------|------|---------|
| ******* ** | A4455988 A4456147 A4456289 A4456311 | 1184 | butyrobetaine (gamma), 2-oxoglutarate | | | |
| ******** ** | A4455988 A4456055 A4456147 A4456289 A4456311 | 1184 | | | | |
| 444 6 44 44 | 44456055 4A456147 4A456289 4A456311 | - | dioxygenase (gamma-butyrobetaine hydroxylase) | 15.54 | down | 0.00001 |
| 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 | AA456147 AA456289 AA456311 | 1185 | EST | 5.16 | down | 0.00158 |
| # # # # # # # # # # # # # # # # # # # | 4A456289 4A456311 | 1188 | general transcription factor IIIA | 4.23 | down | 0.00088 |
| to the state of th | AA456311 | 1189 | EST | 15.31 | down | 0.00004 |
| | | 1190 | EST | 46.81 | down | 0.001 |
| 'at 'at 'at ' | 4A456326 | 1191 | EST | 3.35 | down | 0.00489 |
| , at at at a | AA456589 | 1194 | EST | 4.23 | down | 0.00102 |
| ,a ,a | AA456687 | 1197 | EST | 3.08 | down | 0.01189 |
| , at | AA457377 | 1201 | EST | 3.1 | down | 0.00549 |
| | AA458652 | 1202 | EST | 8.26 | down | 0.00001 |
| rc_AA458923_at A | AA458923 | 1207 | EST | 3.36 | down | 0.00421 |
| at | AA458946 | 1209 | EST | 15.88 | down | 0.00004 |
| at | AA459256 | 1212 | lectin, mannose-binding, 1 | 3.01 | down | 0.00094 |
| at , | AA459293 | 1213 | EST | 3.2 | down | 0.0001 |
| ˈat | AA459389 | 1216 | tyrosylprotein sulfotransferase 2 | 3.72 | down | 0.02252 |
| at | AA459420 | 1217 | EST | 7.25 | down | 0.0214 |
| | AA459668 | 1219 | 3-hydroxyisobutyryl-Coenzyme A hydrolase | 7.62 | down | 0.00225 |
| # | AA459690 | 1221 | EST | 9.18 | down | 0.00732 |
| 1 | | | solute carrier family 22 (extraneuronal | | | |
| at | AA460012 | 1224 | monoamine transporter), member 3 | 4.27 | down | 0.04975 |
| AA460047_at A | AA460047 | 1226 | EST | 3.33 | down | 0.04011 |
| | AA460128 | 1227 | similar to S. pombe dim1+ | 3.28 | down | 0.01299 |
| rc AA460449 at | AA460449 | 1228 | EST | 7.77 | down | 0.00011 |
| at | AA460661 | 1229 | EST | 7.02 | down | 0.00053 |
| at | AA460916 | 1233 | EST | 3.69 | down | 0.04841 |
| at | AA461057 | 1234 | nuclear localization signal deleted in | 5.22 | down | 0.00051 |
| at | AA461303 | 1238 | DKFZP586D1519 protein | 4.77 | down | 0.0438 |
| , at | AA461444 | 1239 | EST | 11.56 | down | 0.00167 |
| ˈatˈ | AA461458 | 1241 | EST | 3.37 | down | 0.02427 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| rc_AA463194_s_at | AA463194 | 1244 | KIAA1037 protein | 4.92 | down | 0.01866 |
| rc AA463195 at | AA463195 | 1245 | EST | 3.41 | down | 0.00413 |
| AA463311_at | AA463311 | 1248 | EST | 3.71 | down | 0.04902 |
| rc AA463729 at | AA463729 | 1250 | EST | 4.07 | down | 0.00676 |
| rc AA463876 at | AA463876 | 1252 | EST | 3.31 | down | 0.00109 |
| rc_AA463946_at | AA463946 | 1254 | pigment epithelium-derived factor | 3.38 | down | 0.0018 |
| rc AA464188 s at | AA464188 | 1256 | EST | 4.82 | down | 0.03208 |
| rc_AA464603_at | AA464603 | 1260 | EST | 3.26 | down | 0.0007 |
| rc_AA465240_at | AA465240 | 1270 | EST | 4.03 | down | 0.0046 |
| rc_AA470153_at | AA470153 | 1275 | solute carrier family 21 (organic anion | 13.26 | down | 0.00315 |
| rc_AA476324_s_at | AA476324 | 1281 | EST | 55.22 | uwop | 0.00132 |
| rc_AA476346_at | AA476346 | 1283 | EST | 3.12 | down | 0.01067 |
| rc_AA476352_at | AA476352 | 1284 | EST | 3.41 | down | 0.02233 |
| rc_AA477119_at | AA477119 | 1289 | EST | 3.13 | down | 0.0338 |
| AA477919_at | AA477919 | 1293 | EST | 4.69 | down | 0.00141 |
| AA477978_s_at | AA477978 | 1294 | short-chain dehydrogenase/reductase 1 | 8.53 | down | 0.01651 |
| rc_AA478416_at | AA478416 | 1300 | EST | 4.04 | down | 0.00078 |
| rc_AA478441_at | AA478441 | 1302 | cathepsin F | 2.07 | down | 0.00752 |
| AA479132_at | AA479132 | 1309 | EST | 3.12 | down | 0.00876 |
| rc_AA479148_at | AA479148 | 1311 | EST | 38.05 | down | 0 |
| rc_AA479488_at | AA479488 | 1313 | S-adenosyfhomocysteine hydrolase-like 1 | 4 | down | 0.0269 |
| rc_AA479498_at | AA479498 | 1314 | EST | 5.78 | down | 0.01489 |
| rc_AA479885_at | AA479885 | 1318 | KIAA0843 protein | 15.57 | down | 0.00024 |
| rc_AA479968_s_at | AA479968 | 1321 | arylsulfatase A | 9.01 | down | 0.00224 |
| rc_AA480975_at | AA480975 | 1322 | EST | 8.95 | down | 0.00259 |
| rc_AA480991_s_at | AA480991 | 1323 | EST · | 8.59 | down | 0.00156 |
| rc_AA481432_s_at | AA481432 | 1328 | fibronectin 1 | 7.76 | down | 0.0061 |
| rc_AA481526_at | AA481526 | 1329 | EST | 3.73 | down | 0.00002 |
| AA481670_at | AA481670 | 1330 | retinal short-chain dehydrogenase/reductase | 6.2 | down | 0.0078 |
| rc_AA482594_at | AA482594 | 1337 | EST | 5.42 | down | 0.00387 |
| rc_AA485089_at | AA485089 | 1341 | EST | 5.46 | down | 0.00044 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| rc_AA485326_at | AA485326 | 1342 | ATP-binding cassette, sub-family D (ALD), | 3.08 | down | 0.00415 |
| rc_AA485413_at | AA485413 | 1344 | EST | 4.54 | down | 0.00137 |
| | | | aldehyde dehydrogenase 5 family, member A1 | | | • |
| rc_AA486410_at | AA486410 | 1348 | (succinate-semialdehyde dehydrogenase) | 8.08 | down | 0.00485 |
| AA486511 at | AA486511 | 1349 | EST | 22.21 | down | 0.00113 |
| rc AA486567 at | AA486567 | 1350 | EST | 2 | down | 0.00002 |
| rc_AA487161_at | AA487161 | 1353 | ubiquilin 2 | 3.13 | down | 0.00023 |
| rc_AA487503_at | AA487503 | 1356 | EST | 8.85 | down | 0.00012 |
| rc_AA487606_at | AA487606 | 1358 | EST | 3.05 | down | 0.00291 |
| rc_AA488843_at | AA488843 | 1362 | cornichon-like | 8.58 | down | 0.02131 |
| rc_AA489061_at | AA489061 | 1367 | EST | 3.49 | down | 0.00223 |
| rc_AA489629_at | AA489629 | 1369 | EST | 8.08 | down | 0.00109 |
| rc_AA489636_at | AA489636 | 1370 | EST | 10.7 | down | 0 |
| rc_AA489798_at | AA489798 | 1373 | hypothetical protein, estradiol-induced | 8.75 | down | 0.00544 |
| rc AA490159 at | AA490159 | 1374 | glucose-6-phosphatase, transport (glucose-6- | 5.44 | down | 0 |
| rc_AA490214_at | AA490214 | 1376 | EST | 3.12 | down | 0.02382 |
| rc_AA490620_at | AA490620 | 1378 | EST | 4.77 | down | 0.00201 |
| rc_AA490670_at | AA490670 | 1379 | EST | 96.6 | down | 0.00454 |
| | | | UDP-N-acetylglucosamine-2-epimerase/N- | | | |
| AA490775_at | AA490775 | 1380 | acetylmannosamine kinase | 5.34 | down | 0.00118 |
| rc AA490882 s at | AA490882 | 1381 | EST | 3.29 | down | 0.00319 |
| rc_AA490890_at | AA490890 | 1382 | EST | 3.02 | down | 0.00007 |
| rc_AA491000_at | AA491000 | 1385 | EST | 4.23 | down | 0.02305 |
| rc_AA491001_i_at | AA491001 | 1386 | EST | 8.52 | down | 0.01118 |
| rc AA491001 f at | AA491001 | 1386 | EST | 3.73 | down | 0.01957 |
| AA495758 s at | AA495758 | 1391 | EST | 3.94 | uwop | 0.00772 |
| rc_AA495820_at | AA495820 | 1393 | EST | 3.98 | down | 0.00218 |
| rc_AA496053_at | AA496053 | 1396 | EST | 3.28 | down | 0.00095 |
| AA496423 at | AA496423 | 1399 | WW domain binding protein 2 | 3.52 | down | 0.01314 |
| rc_AA496914_at | AA496914 | 1401 | v-maf musculoaponeurotic fibrosarcoma (avian) | 3.48 | down | 0.00361 |
| rc_AA497052_at | AA497052 | 1408 | DKFZP727G051 protein | 7.28 | down | 0.01745 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| rc_AA504492_at | AA504492 | 1414 | tubulin, alpha, ubiquitous | 4.21 | down | 0.00752 |
| AA505198 at | AA505198 | 1419 | EST | 3.41 | down | 0.0343 |
| rc AA521290 at | AA521290 | 1421 | EST | 4.53 | down | 0.0148 |
| rc_AA521292_at | AA521292 | 1422 | EST | 8.58 | down | 0.00064 |
| rc_AA521306_at | AA521306 | 1423 | EST | 4.27 | down | 0.00567 |
| 1 | | | UDP-N-acetylglucosamine-2-epimerase/N- | | | |
| rc_AA598417_at | AA598417 | 1426 | acetylmannosamine kinase | 8.56 | down | 0.01638 |
| rc_AA598419_s_at | AA598419 | 1427 | translational inhibitor protein p14.5 | 23.6 | down | 0.00036 |
| rc AA598675 at | AA598675 | 1433 | EST | 3.25 | down | 0.03934 |
| rc_AA598679_at | AA598679 | 1434 | EST | 5.37 | down | 0.00467 |
| rc AA598746 at | AA598746 | 1437 | EST | 3.8 | down | 0.02667 |
| rc_AA598926 at | AA598926 | 1441 | EST | 3.7 | down | 0.00432 |
| rc_AA599211_at | AA599211 | 1445 | short-chain dehydrogenase/reductase 1 | 7.85 | down | 0.00911 |
| rc_AA599234_s_at | AA599234 | 1447 | murine leukemia viral (bmi-1) oncogene homolog | 3.9 | down | 0.0068 |
| rc_AA599472_at | AA599472 | 1451 | succinate-CoA ligase, GDP-forming, beta subunit | 5.07 | down | 0.00447 |
| rc_AA599526_at | AA599526 | 1453 | cartilage associated protein | 3.02 | down | 0.00043 |
| rc_AA599814_at | AA599814 | 1456 | EST | 12.37 | down | 0.00002 |
| rc_AA599937_s_at | AA599937 | 1458 | insulin-like growth factor-binding protein 4 | 26.92 | down | 0.00094 |
| rc_AA599954_at | AA599954 | 1459 | cell cycle progression 8 protein | 3.15 | down | 0.00021 |
| rc_AA608546_at | AA608546 | 1463 | EST | 12.52 | down | 0.00003 |
| rc_AA608671_at | AA608671 | 1466 | EST | 3.14 | down | 0.04543 |
| rc_AA608729_at | AA608729 | 1468 | EST | 3.89 | down | 0.01757 |
| rc_AA608751_i_at | AA608751 | 1469 | EST | 5.76 | down | 0.01404 |
| rc_AA608802_at | AA608802 | 1470 | EST | 6.95 | down | 0.00263 |
| rc_AA608807_s_at | AA608807 | 1471 | inhibin, beta B (activin AB beta polypeptide) | 4.05 | down | 0.00568 |
| rc_AA608837_at | AA608837 | 1472 | EST | 6.2 | down | 0.00006 |
| rc_AA609011_at | AA609011 | 1476 | EST | 3.94 | down | 0.0313 |
| rc_AA609164_at | AA609164 | 1480 | cytochrome b-561 | 6.8 | down | 0.02298 |
| rc_AA609316_at | AA609316 | 1481 | EGF-like-domain, multiple 5 | 7.97 | down | 0.00011 |
| rc_AA609519_at | AA609519 | 1482 | EST | 8.13 | down | 600000 |
| rc_AA609537_s_at | AA609537 | 1483 | hepatic leukemia factor | 8.76 | down | 0.00018 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seg ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| rc_AA609572_at | AA609572 | 1484 | EST | 3.02 | down | 0.01534 |
| rc AA609574 at | AA609574 | 1485 | EST | 5.03 | down | 0 |
| rc_AA609576_at | AA609576 | 1486 | EST | 3.1 | down | 0.00266 |
| rc_AA609773_at | AA609773 | 1489 | EST | 60.9 | down | 0.01103 |
| rc_AA609774_at | AA609774 | 1490 | EST | 4.02 | down | 0.00424 |
| rc_AA609934_at | AA609934 | 1493 | EST | 6.84 | down | 0.00048 |
| rc_AA609996_at | AA609996 | 1495 | EST | 3.93 | down | 0.00988 |
| rc AA620343 at | AA620343 | 1500 | EST | 5.04 | down | 0.00407 |
| rc_AA620556_at | AA620556 | 1505 | EST | 32.4 | down | 0.00353 |
| rc_AA620667_s_at | AA620667 | 1506 | protein tyrosine phosphatase type IVA, member 1 | 5.92 | down | 0.00206 |
| rc_AA620830_at | AA620830 | 1509 | DKFZP5641122 protein | 3.42 | down | 0.02421 |
| rc_AA621131_at | AA621131 | 1513 | EST | 35.37 | down | 0 |
| rc_AA621192_at | AA621192 | 1515 | EST | 5.39 | down | 0.0016 |
| rc_AA621209_at | AA621209 | 1516 | similar to Caenorhabditis elegans protein | 6.34 | down | 0.00144 |
| rc_AA621235_at | AA621235 | 1517 | EST | 3.44 | down | 0.0021 |
| rc_AA621274_i_at | AA621274 | 1519 | EST | 7.43 | down | 0.00065 |
| rc_AA621430_at | AA621430 | 1525 | doublecortex; lissencephaly, X-linked | 3.09 | down | 0.00024 |
| rc_AA621796_at | AA621796 | 1531 | kinesin family member 3B | 4.44 | down | 0.00032 |
| | | | PDZ domain containing guanine nucleotide | | | |
| AB002311_at | AB002311 | 1535 | exchange factor(GEF)1; RA(Ras/Rap1A- | 4.21 | down | 0.00476 |
| AB002328_at | AB002328 | 1536 | calcineurin binding protein 1 | 5.55 | down | 0.00016 |
| AF000573_rna1_at | AF000573 | 1543 | homogentisate 1,2-dioxygenase (homogentisate | 13.76 | down | 0.00002 |
| AF005039_at | AF005039 | 1548 | secretory carrier membrane protein 3 | 3.42 | down | 0.04953 |
| | | | solute carrier family 4, sodium bicarbonate | | | |
| AF007216_at | AF007216 | 1550 | cotransporter, member 4 | 5.79 | down | 0.00005 |
| C01257_at | C01257 | 1554 | EST | 5.35 | down | 0.00608 |
| C01286_s_at | C01286 | 1555 | integral membrane protein 2B | 4.11 | down | 0.00292 |
| C01409_s_at | C01409 | 1556 | EST | 4.41 | down | 0.01725 |
| C01686_at | C01686 | 1557 | EST | 3.01 | down | 0.00048 |
| C02099 s at | C02099 | 1560 | CGI-131 protein | 5.85 | down | 0.02377 |
| C02460_at | C02460 | 1562 | EST | 3.64 | down | 0.02705 |
| | | | | | | |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| rc C14963 s at | C14963 | 1572 | nicotinamide nucleotide transhydrogenase | 3.9 | down | 0.0044 |
| C15871 at | C15871 | 1575 | EST | 3.26 | down | 0.00046 |
| C16420 s at | C16420 | 1576 | EST. | 5.95 | down | 0.00119 |
| rc C20653 at | C20653 | 1578 | EST | 10.59 | down | 0.00001 |
| rc_C20810_at | C20810 | 1579 | EST | 5.17 | down | 0.00614 |
| rc_C20911_at | C20911 | 1580 | antithrombin III | 6.56 | down | 0.00175 |
| rc_C20974_at | C20974 | 1581 | Vanin 1 | 99.9 | down | 0.00272 |
| rc_C21130_at | C21130 | 1583 | EST | 8.79 | down | 0.00008 |
| rc C21238 at | C21238 | 1584 | EST | 4.54 | down | 0.02074 |
| 1 | | • | cytochrome P450, subfamily IIIA (niphedipine | | | |
| | D00003 | 1586 | oxidase), polypeptide 3 | 22.05 | down | 0.00059 |
| ! | | | cytochrome P450, subfamily IIIA (niphedipine | | | |
| D00003 at | D00003 | 1586 | oxidase), polypeptide 3 | 9.46 | down | 0.00001 |
| D00097 s at | D00097 | 1588 | amyloid P component, serum | 16.72 | down | 0.00098 |
| i I | | | cytochrome P450, subfamily IIIA (niphedipine | | | |
| | | | oxidase), polypeptide 3, cytochrome P450, | | | |
| | | | subfamily IIIA (niphedipine oxidase), polypeptide | | | |
| D00408 s at | D00408 | 1589 | 5, cytochrome P450, subfamily IIIA, polypeptide 7 | 11.1 | down | 0 |
| D00632 at | D00632 | 1591 | glutathione peroxidase 3 (plasma) | 6.55 | down | 0.00121 |
| D00723_at | D00723 | 1592 | glycine cleavage system protein H (aminomethyl | 4.18 | down | 0.00543 |
| D10040_at | D10040 | 1593 | fatty-acid-Coenzyme A ligase, long-chain 2 | 20.51 | down | 0 |
| | | | acetyl-Coenzyme A acetyltransferase 1 | | | |
| D10511_at | D10511 | 1594 | (acetoacetyl Coenzyme A thiolase) | 10.68 | down | 0.0002 |
| rc D11756 f at | D11756 | 1596 | EST | 5.49 | down | 0.01272 |
| rc D11802 at | D11802 | 1597 | angiotensinogen | 5.65 | down | 0.0000 |
| rc_D11835_at | D11835 | 1598 | low density lipoprotein receptor (familial | 21.76 | down | 0.00307 |
| rc_D11881_at | D11881 | 1599 | KIAA0962 protein | 4.37 | down | 0.01627 |
| 1 : | 1 | , | | 1 | 9 | 00000 |
| D12485_at | D12485 | 1600 | 1 (nomologous to mouse Ly-41 antigen) | 4.5/ | down | 0.0000 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change Direction | Direction | Pvalue |
|--------------|---------|--------|--|-----------------------|-----------|---------|
| | | | cytochrome P450, subfamily IVF, polypeptide 2, cytochrome P450, subfamily IVF, polypeptide 3 | | : | |
| D12620_s_at | D12620 | 1601 | (leukotriene B4 omega hydroxylase) cytochrome P450, subfamily IVF, polypeptide | 35.09 | down | 0.00015 |
| D12620 s at | D12620 | 1601 | 2,cytochrome P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) | 13.45 | down | C |
| D13243 s at | D13243 | 1602 | pyruvate kinase, liver and RBC | 20.22 | down | 0 |
| D13643_at | D13643 | 1609 | KIAA0018 gene product | 10.84 | down | 0.00058 |
| D13705_s_at | D13705 | 1610 | cytochrome P450, subfamily IVA, polypeptide 11 | 3.7 | down | 0.00038 |
| D13814_s_at | D13814 | 1611 | angiotensin receptor 1, angiotensin receptor 1B | 3.12 | down | 0.00101 |
| D14012_s_at | D14012 | 1612 | HGF activator | 12.75 | down | 0.0035 |
| D14664_at | D14664 | 1616 | KIAA0022 gene product | 8.98 | down | 0.00011 |
| D14695_at | D14695 | 1618 | KIAA0025 gene product; MMS-inducible gene | 6.48 | down | 0 |
| | | | acetyl-Coenzyme A acyltransferase 2 | | | |
| D16294_at | D16294 | 1619 | (mitochondrial 3-oxoacyl-Coenzyme A thiolase) | 4.81 | down | 0.03921 |
| D16350_at | D16350 | 1620 | SA (rat hypertension-associated) homolog | 3.83 | down | 0.00117 |
| | | | hydroxyacyl-Coenzyme A dehydrogenase/3- | | | |
| | | | ketoacyl-Coenzyme A thiolase/enoyi-Coenzyme | | | |
| D16481_at | D16481 | 1621 | A hydratase (trifunctional protein), beta subunit | 3.18 | down | 0.00695 |
| D16626_at | D16626 | 1622 | histidine ammonia-lyase | 22.66 | down | 0 |
| D16626_at | D16626 | 1622 | histidine ammonia-lyase | 9.25 | down | 0.00025 |
| rc_D20350_át | D20350 | 1624 | EST | 13.97 | down | 0.00057 |
| D31117_at | D31117 | 1640 | ribosome binding protein 1 (dog 180kD homolog) | 5.3 | down | 0.02749 |
| D31225_at | D31225 | 1641 | EST | 3.17 | down | 0.01073 |
| D31289_at | D31289 | 1642 | EST | 4.16 | down | 0.02166 |
| D31381_at | D31381 | 1644 | dynein, axonemal, light polypeptide 4 | 4.97 | down | 0.01806 |
| D31628_s_at | D31628 | 1646 | 4-hydroxyphenylpyruvate dioxygenase | 50.48 | down | 0.00002 |
| D31716_at | D31716 | 1647 | basic transcription element binding protein 1 | 5.35 | down | 0.00086 |
| D31815_at | D31815 | 1648 | regucalcin (senescence marker protein-30) | 10.55 | down | 0.00037 |
| D31887_at | D31887 | 1649 | KIAA0062 protein | 4.26 | down | 0.00101 |
| D37931_at | D37931 | 1650 | ribonuclease, RNase A family, 4 | 5.81 | down | 0.00836 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction Pvalue | Pvalue |
|----------------|---------|--------|---|-------------|------------------|---------|
| | | | inter-alpha (globulin) inhibitor H4 (plasma | | | |
| D38535_at | D38535 | 1654 | Kallikrein-sensitive glycoprotein) | 13.2 | down | 0.01165 |
| rc_D45529_at | D45529 | 1662 | EST | 3.82 | down | 0.00193 |
| rc_D45556_at | D45556 | 1663 | EST | 4.83 | down | 0.01044 |
| rc D45714 at | D45714 | 1664 | EST | 5.64 | down | 0.00384 |
| D49357 at | D49357 | 1665 | methionine adenosyltransferase I, alpha | 11.28 | down | 0.00331 |
| D49387_at | D49387 | 1666 | NADP dependent leukotriene b4 12- | 8.17 | down | 0.00972 |
| D49742_at | D49742 | 1668 | hyaluronan-binding protein 2 | 18.13 | down | 0.00012 |
| rc_D51199_at | D51199 | 1677 | EST | 5.05 | down | 0.00192 |
| rc_D51279_s_at | D51279 | 1679 | ovarian granulosa cell protein (13kD) | 5.88 | down | 0.01271 |
| rc_D52097_s_at | D52097 | 1682 | prostatic binding protein | 8.1 | down | 0.00141 |
| D57823_at | D57823 | 1690 | Sec23 (S. cerevisiae) homolog A | 4.43 | down | 0 |
| D58231_s_at | D58231 | 1692 | ubiquitin-like 3 | 3.07 | down | 0.0002 |
| rc_D59344_s_at | D59344 | 1695 | EST | 3.34 | down | 0.01337 |
| rc_D59554_f_at | D59554 | 1698 | EST | 6.7 | down | 0 |
| rc_D59714_s_at | D59714 | 1700 | mitogen inducible 2 | 17.62 | down | 0.00014 |
| rc_D60670_at | De0670 | 1702 | EST | 3.73 | down | 0.00382 |
| rc_D60769_s_at | D60769 | 1703 | KIAA0096 protein | 4.31 | down | 0.00142 |
| rc_D60856_f_at | De0856 | 1705 | UDP-glucose dehydrogenase | 6.45 | down | 0.01222 |
| D61991_at | D61991 | 1706 | EST | 4.84 | down | 0.00005 |
| D62103_s_at | D62103 | 1707 | EST | 4.11 | down | 0.0263 |
| rc_D62518_at | D62518 | 1708 | EST | 17.49 | down | 0.00017 |
| D63160_at | D63160 | 1709 | ficolin (collagen/fibrinogen domain-containing | 4.01 | down | 0.00391 |
| D78011_at | D78011 | 1717 | dihydropyrimidinase | 21.37 | down | 0.00003 |
| D79276_at | D79276 | 1722 | succinate-CoA ligase, GDP-forming, beta subunit | 8.9 | down | 0.00047 |
| D79687_at | D79687 | 1723 | KIAA1053 protein | 5.06 | down | 0.00047 |
| rc_D80050_at | D80050 | 1726 | EST | 4.64 | down | 0.01001 |
| rc_D80217_f_at | D80217 | 1727 | H91620p protein | 3.61 | down | 0.01973 |
| rc_D80218_f_at | D80218 | 1728 | brain acid-soluble protein 1 | 3.83 | down | 0.0137 |
| rc_D80312_f_at | D80312 | 1730 | EST | 3.74 | down | 0.01909 |
| rc_D80408_at | D80408 | 1731 | EST | 3.36 | down | 0.00102 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affv ID | Genbank | Sea ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| rc_D80905_at | D80905 | 1735 | EST | 3.59 | down | 0.0007 |
| D82061_at | D82061 | 1741 | Ke6 gene, mouse, human homolog of | 6.1 | down | 0.00104 |
| D82422_at | D82422 | 1745 | ferroportin 1; iron regulated gene 1 | 6.01 | down | 0.02351 |
| D85181_at | D85181 | 1750 | sterol-C5-desaturase (fungal ERG3, delta-5- | 9.56 | down | 0.00005 |
| D86062_s_at | D86062 | 1752 | ES1 (zebrafish) protein, human homolog of | 3.63 | down | 0.0001 |
| D87075_at | D87075 | 1760 | solute carrier family 23 (nucleobase transporters), | 4.15 | down | 0.00067 |
| D87436_at | D87436 | 1761 | KIAA0249 gene product | 5.49 | down | 0.00333 |
| D87449_at | D87449 | 1762 | KIAA0260 protein | 4.58 | down | 0.00026 |
| D87466_at | D87466 | 1763 | KIAA0276 protein | 4.83 | down | 0.0007 |
| D90042_at | D90042 | 1767 | N-acetyltransferase 2 (arylamine N- | 7.06 | down | 0 |
| D90282_at | D90282 | 1769 | carbamoyl-phosphate synthetase 1, mitochondrial | 27.29 | down | 0.00002 |
| rc_F02028_at | F02028 | 1774 | EST | 23.48 | down | 0.00465 |
| rc_F02094_at | F02094 | 1775 | ecotropic viral integration site 5 | 3.41 | down | 0.00495 |
| rc_F02245_at | F02245 | 1776 | monoamine oxidase A | 3.9 | down | 0.02943 |
| rc_F02345_at | F02345 | 1779 | EST | 3.9 | down | 0.0033 |
| rc_F03200_at | F03200 | 1783 | EST | 3.75 | down | 0.01805 |
| | | | matrix metalloproteinase 2 (gelatinase A, 72kD | | | |
| rc_F03969_at | F03969 | 1785 | gelatinase, 72kD type IV collagenase) | 7.87 | down | 0.00014 |
| rc_F04335_at | F04335 | 1787 | EST | 3.16 | down | 0.0058 |
| rc_F04611_at | F04611 | 1792 | EST | 23.96 | down | 0.00018 |
| rc_F04944_s_at | F04944 | 1795 | acyl-Coenzyme A oxidase | 4.01 | down | 0.00242 |
| rc_F08817_at | F08817 | 1796 | EST | 8.29 | down | 0.0077 |
| rc_F08941_at | F08941 | 1798 | EST | 3.48 | down | 0.00428 |
| rc_F09058_at | F09058 | 1799 | EST | 3.6 | down | 0.00595 |
| rc_F09350_at | F09350 | 1801 | EST | 4.79 | down | 0.00088 |
| rc_F09353_at | F09353 | 1802 | solute carrier family 5 (inositol transporters), | 3.3 | down | 0.02841 |
| rc F09578 at | F09578 | 1804 | translocated to. 3 | 4.66 | down | 0.04463 |
| rc_F09979_at | F09979 | 1809 | EST | 4.36 | down | 0.02555 |
| rc F10182 s at | F10182 | 1812 | hepsin (transmembrane protease, serine 1) | 58.92 | down | 0.00837 |
| rc_F10276_s_at | F10276 | 1814 | dual specificity phosphatase 6 | 8.13 | down | 0.0001 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| rc_F10466_at rc_F10640_at rc_F10874_f_at rc_F10875_at rc_F13702_at rc_F13782_s_at | | | | 5 20 | איייני | 0.02494 |
|---|-----------|-----------|--|-------|--------|---------|
| rc_F10640_at rc_F10874_f_at rc_F10875_at rc_F13702_at rc_F13782_s_at | | | | 25.0 | | 101100 |
| rc_F10874_f_at rc_F10875_at rc_F13702_at rc_F13782_s_at | | 1821 EST | | 3.58 | down | 0.00152 |
| rc_F10875_at rc_F13702_at rc_F13782_s_at | 10874 18 | 1823 EST | - | 4.19 | down | 0.00025 |
| rc_F13702_at rc_F13782_s_at | , | 1824 EST | h- | 5.09 | down | 0.00004 |
| rc_F13782_s_at | 13702 18 | 1826 EST | — | 6.01 | down | 0.00064 |
| | 13782 18 | 1827 LIIV | _IM binding domain 2 | 4.17 | down | 0.00109 |
| | | 田田 | IEK tyrosine kinase, endothelial (venous | | | |
| rc_H02848_s_at | 102848 18 | 1831 ma | malformations, multiple cutaneous and mucosal) | 3.47 | down | 0.0089 |
| rc_H02855_at | 102855 18 | 1832 EST | L | 5.96 | down | 0.00458 |
| rc_H03348_at · · · · | 103348 18 | 1833 claı | claudin 1 | 5.77 | down | 0.0001 |
| rc_H03945_at | 103945 18 | 1835 EST | L | 4.94 | down | 0.02603 |
| rc_H04142_f_at h | 104142 18 | 1836 EST | ļ. | 3.57 | down | 0.01906 |
| rc_H04242_at | 104242 18 | 1837 RA | RAB5B, member RAS oncogene family | 3.27 | down | 0.04826 |
| H04854_at H | 104854 18 | 1842 inte | nterleukin 1 receptor accessory protein | 6.58 | down | 0.00007 |
| rc_H05072_at | 105072 18 | 1843 EST | — | 3.12 | down | 0.01248 |
| rc_H05974_s_at | 105974 18 | 1850 EST | F | 6.28 | down | 0.00549 |
| rc_H05985_at H | 105985 18 | _ | hypothetical protein | 6.43 | down | 0.04887 |
| rc_H06063_s_at | 106063 18 | | chondroitin sulfate proteoglycan 3 (neurocan) | 3.15 | down | 0.00599 |
| rc_H06144_at | 106144 18 | 1853 EST | | 3.1 | nwob. | 0.00745 |
| rc_H06166_at | 106166 18 | 1854 EST | - | 3.31 | down | 0.03778 |
| rc_H06935_s_at | 106935 18 | 1855 ele | electron-transferring-flavoprotein dehydrogenase | 6.82 | down | 0.00105 |
| . rc_H08054_at | 108054 18 | 857 EST | | 4.2 | down | 6000.0 |
| rc_H08102_at | 108102 18 | 1858 bre | breast cell glutaminase | 27.77 | down | 0.00032 |
| rc_H09167_at | 109167 18 | 1860 KIA | KIAA0195 gene product | 3.31 | down | 0.00313 |
| rc_H09353_at | 109353 18 | 1866 EST | | 23.06 | down | 0.00094 |
| H09364_s_at | 109364 18 | 867 suc | succinate dehydrogenase complex, subunit A, | 5.74 | down | 0.03125 |
| rc_H09594_at | 109594 18 | 1868 EST | — | 3.12 | down | 0.00231 |
| rc_H09959_s_at | H09959 18 | 869 cho | choline kinase | 3.25 | down | 0.00225 |
| H10482_at | 110482 18 | 1870 EST | — | 3.19 | down | 0.01611 |
| rc_H10661_at | 110661 18 | 1871 EST | . | 4.54 | down | 0.00276 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction . Pvalue | Pvalue |
|----------------|---------|--------|---|-------------|--------------------|---------|
| | | | methylenetetrahydrofolate dehydrogenase (NADP+ dependent) methenytetrahydrofolate | | - | |
| rc H10779 s at | H10779 | 1872 | cyclohydrolase, formylletrahydrofolate synthetase | 9 73 | down | 0.00035 |
| rc H11274 at | H11274 | 1874 | EST | 4.13 | down | 0.01478 |
| | • | | glycine dehydrogenase (decarboxylating; glycine | | | |
| rc H11739 s at | H11739 | 1876 | decarboxylase, glycine cleavage system protein | 10.33 | down | 0.00023 |
| rc H11746 at | H11746 | 1877 | EST | 3.92 | down | 0.00012 |
| rc_H12257_at | H12257 | 1879 | EST | 3.19 | down | 6900.0 |
| rc_H12593_at | H12593 | 1880 | zinc-finger protein 265 | 10.72 | down | 0.0056 |
| rc_H13696_at | H13696 | 1882 | EST | 3.48 | down | 0.01796 |
| rc H14372 s at | H14372 | 1883 | ATP-binding cassette, sub-family A (ABC1), | 5.16 | down | 0.00012 |
| rc_H16768_at | H16768 | 1887 | EST | 3.72 | down | 0.00688 |
| rc_H18950_at | H18950 | 1892 | EST | 3.85 | down | 0.00162 |
| rc_H18997_at | H18997 | 1893 | F-box protein 21 | 3.87 | down | 0.00611 |
| rc H19504 f at | H19504 | 1895 | EST | 3.13 | down | 0.04948 |
| rc_H20543_at | H20543 | 1897 | DKFZP586B1621 protein | 31.03 | down | 0.00074 |
| rc_H25124_at | H25124 | 1903 | EST | 3.65 | down | 0.00004 |
| rc_H25551_at | H25551 | 1904 | EST | 3.54 | down | 0.00366 |
| rc_H25836_at | H25836 | 1905 | tumor necrosis factor (ligand) superfamily, | 3.3 | down | 0.03125 |
| rc_H26417_at | H26417 | 1906 | EST | 3.22 | down | 0.03672 |
| rc_H26763_at | H26763 | 1907 | EST | 3.39 | down | 0.04188 |
| rc_H27330_at | H27330 | 1909 | EST | 3.2 | down | 0.00067 |
| rc H27442 s_at | H27442 | 1910 | erythrocyte membrane protein band 7.2 | 6.81 | down | 0.00083 |
| rc_H29568_at | H29568 | 1914 | EST | 11.45 | down | 0.00058 |
| rc_H30270_at | H30270 | 1915 | EST | 17.09 | down | 0.00001 |
| rc_H38246_s_at | H38246 | 1917 | EST | 9.25 | down | 0.00157 |
| rc_H39119_at | H39119 | 1919 | EST | 3.06 | down | 0.03349 |
| rc_H40149_at | H40149 | 1921 | KIAA0937 protein | 4.59 | down | 0.00112 |
| rc_H40424_s_at | H40424 | 1922 | butyrate response factor 1 (EGF-response factor | 3.56 | down | 0.04066 |
| rc_H40534_at | H40534 | 1923 | EST | 3.18 | down | 0.01381 |
| rc_H41084_at | H41084 | 1924 | EST | 6.31 | down | 0.0227 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| rc H41280 at | H41280 | 1925 | EST | 3.68 | down | 0.00455 |
| rc H42053 s at | H42053 | 1927 | EST | 3.49 | down | 0.01057 |
| rc H46001 at | H46001 | 1931 | EST | 5.03 | down | 0.00563 |
| H46990 at | H46990 | 1933 | cytochrome P450, subfamily IIE (ethanol- | 3.2 | down | 0.00095 |
| rc H47391 at | H47391 | 1935 | EST | 3.1 | down | 0.03807 |
| rc_H47838_at | H47838 | 1936 | carboxypeptidase B2 (plasma) | 16.74 | down | 0.00002 |
| rc_H49415_at | H49415 | 1938 | EST | 3.72 | down | 0.0005 |
| H51340 at | H51340 | 1941 | EST | 3.73 | down | 0.02643 |
| rc H54285 s at | H54285 | 1947 | EST | 5.14 | down | 0.00426 |
| rc H55759 at | H55759 | 1949 | EST | 11.52 | down | 0.00034 |
| 1 | | | 4-nitrophenylphosphatase domain and non- | | | |
| rc H56584 at | H56584 | 1951 | neuronal SNAP25-like 1 | 9.5 | down | 0 |
| rc H57060 s at | H57060 | 1954 | EST | 30.98 | down | 0.01687 |
| rc H57166 at | H57166 | 1955 | EST | 60.76 | down | 0.00007 |
| rc_H57816_at | H57816 | 1957 | EST | 4.41 | down | 0.00206 |
|]] | | | protein phosphatase 2 (formerly 2A), regulatory | | | |
| rc_H57850_at | H57850 | 1958 | subunit A (PR 65), beta isoform | 3.02 | down | 0.00123 |
| rc_H58673_at | H58673 | 1959 | EST | 14.85 | down | 0.00005 |
| rc h58692 s_at | H58692 | 1960 | formyltetrahydrofolate dehydrogenase | 81.41 | down | 0 |
| rc_H59136_at | H59136 | 1962 | EST | 8.64 | down | 0.00013 |
| rc_H59141_at | H59141 | 1963 | EST | 3.12 | down | 0.00293 |
| rc H60595 s at | H60595 | 1966 | progesterone binding protein | 15.8 | down | 0.01078 |
| H61295 s at | H61295 | 1968 | CD4 antigen (p55) | 10.71 | down | 0.00925 |
| rc H62838 at | H62838 | 1971 | EST | 3.09 | down | 0.03201 |
| rc_H63251_at | H63251 | 1972 | KIAA0606 protein; SCN Circadian Oscillatory | 3.27 | down | 0.02455 |
| rc_H65650_at | H65650 | 1976 | EST | 3.88 | down | 0.0083 |
| H66367 at | H66367 | 1977 | EST | 6.68 | down | 0.0001 |
| rc_H66840_at | H66840 | 1978 | EST | 3.67 | down | 0.0143 |
| rc_H67094_at | H67094 | 1979 | EST | 3.24 | down | 0.00075 |
| H67840_at | H67840 | 1980 | EST | 3.1 | down | 0.00528 |
| rc H68097 at | H68097 | 1982 | EST | 3.83 | down | 0.00797 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| H68953_at | H68953 | 1985 | transferrin | 6.4 | | 0.00132 |
| rc_H69138_at | H69138 | 1986 | v-Ki-ras2 Kirsten rat sarcoma 2 viral oncogene | 6.76 | down | 0.00142 |
| rc_H69565_at | H69565 | 1987 | EST | 4.11 | down | 0.00002 |
| rc_H70554_at | H70554 | 1989 | EST | 10.99 | down | 0 |
| rc_H71169_at | H71169 | 1992 | putative protein similar to nessy (Drosophila) | 4 | down | 0.00709 |
| | | | glucan (1,4-alpha-), branching enzyme 1 | | | |
| rc_H71861_s_at | H71861 | 1993 | (glycogen branching enzyme, Andersen disease, | 5.97 | down | 0.00007 |
| rc_H73535_s_at | H73535 | 1996 | EST | 6.83 | down | 0.00202 |
| rc_H74317_s_at | H74317 | 1997 | apolipoprotein A-II | 45.09 | down | 0.01982 |
| ۱-۳ | H77597 | 2000 | metallothionein 1H | 16.03 | down | 0.00675 |
| H78628_at | H78628 | 2003 | EST | 4.98 | down | 0.00729 |
| rc_H79820_at | H79820 | 2004 | EST | 3.25 | down | 0.01466 |
| rc_H80901_s_at | H80901 | 2002 | ficolin (collagen/fibrinogen domain-containing) 3 | 50.61 | down | 0.00262 |
| rc_H81070_f_at | H81070 | 2006 | RNA helicase-related protein | 25.74 | down | 0.00126 |
| rc_H82966_s_at | H82966 | 2011 | apolipoprotein B (including Ag(x) antigen) | 3.42 | down | 0.00769 |
| rc_H83109_f_at | H83109 | 2012 | EST | 16.55 | down | 0.00001 |
| rc_H83442_s_at | H83442 | 2013 | catechol-O-methyltransferase | 3.99 | down | 0.00594 |
| rc_H83451_at | H83451 | 2014 | EST | 3.35 | down | 0.00498 |
| rc_H87144_at | H87144 | 2016 | EST | 3.41 | down | 0.00387 |
| rc_H87765_at | H87765 | 2017 | KIAA0626 gene product | 3.86 | down | 0.00131 |
| H88033_s_at | H88033 | 2019 | KIAA0733 protein | 4.42 | down | 0.02032 |
| rc_H88359_s_at | H88359 | 2020 | nuclear factor (erythroid-derived 2)-like 2 | 5.16 | down | 0.01253 |
| rc_H88675_at | H88675 | 2022 | EST | 5.63 | down | 0.00554 |
| rc_H89514_s_at | H89514 | 2023 | protein kinase, cAMP-dependent, catalytic, alpha | 3.44 | down | 0.00435 |
| rc_H89893_at | H89893 | 2025 | EST | 3.17 | down | 0.00658 |
| rc_H89980_at | H89980 | 2026 | protein phosphatase 1, regulatory (inhibitor) | 31.13 | down | 0.00006 |
| rc_H90417_s_at | H90417 | 2028 | EST | 4.17 | down | 0.015 |
| rc_H91325_s_at | H91325 | 2029 | aldolase B, fructose-bisphosphate | 45.85 | down | 0.00505 |
| rc_H91456_s_at | H91456 | 2030 | nuclear receptor subfamily 1, group H, member 4 | 4.9 | down | 0.00255 |
| | | | phosphorylase, glycogen; liver (Hers disease, | | | |
| rc_H91680_s_at | H91680 | 2032 | glycogen storage disease type VI) | 4.15 | down | 0.00746 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

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|----------|--|-----------------|--|--|
| a Dig | glutamate-cysteine ligase (gamma- | | | |
| 2034 glu | glutamylcysteine synthetase), catalytic (72.8kD) | 5.06 | down | 0.01029 |
| 2035 EST | - | 15.3 | down | 0.00233 |
| 2036 EST | F | 24.23 | down | 0 |
| 2038 pro | proline synthetase co-transcribed (bacterial | 3.17 | down | 0.00113 |
| ले | lydroxy-3-methylglutaryl-Coenzyme A synthase | | | |
| 2041 2 (| mitochondrial) | 9.49 | down | 0.02373 |
| 2043 alp | ha-2-plasmin inhibitor | 40.92 | down | 0.00271 |
| 2044 ES | <u></u> | 4.77 | down | 0.00266 |
| | ha-1-B glycoprotein | 47.03 | down | 0.01158 |
| | L | 3.17 | down | 0.00182 |
| _ | FZP586A0522 protein | 28.48 | down | 0.00139 |
| | | 8.55 | down | 0.00046 |
| | F | 4.02 | down | 0.01565 |
| | <u></u> | 3.86 | down | 0.00362 |
| | <u></u> | 3.86 | down | 0.01534 |
| | | 4.66 | down | 0.03722 |
| | L- | 5.09 | down | 0.00025 |
| | :L2/adenovirus E1B 19kD-interacting protein 3 | 80 | down | 0.0018 |
| | <u> </u> | 3.31 | down | 0.00174 |
| | | 4.38 | down | 0.00548 |
| | <u></u> | 3.57 | down | 0.00298 |
| | dothelin receptor type B | 3.43 | down | 0.00093 |
| | ipose differentiation-related protein; adipophilin | 5.83 | down | 0.04346 |
| | erleukin 6 signal transducer (gp130, oncostatin | 3.59 | down | 0.00366 |
| | | _ · · · · · _ , | proline synthetase co-franscribed (bacterial 3-hydroxy-3-methylglutaryl-Coenzyme A synthase 2 (mitochondrial) alpha-2-plasmin inhibitor EST alpha-1-B glycoprotein EST | proline synthetase co-transcribed (bacterial 3-17 3-hydroxy-3-methylglutaryl-Coenzyme A synthase 2 (mitochondrial) alpha-2-plasmin inhibitor EST 4.77 alpha-1-B glycoprotein 3.17 47.03 EST 5.55 EST EST 8.55 endothelin receptor type B adipose differentiation-related protein; adipophilin 5.83 interleukin 6 signal transducer (gp130, oncostatin 3.55 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|---------------|---------|--------|---|-------------|-----------|---------|
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| | | | | | | |
| J02843 at | J02843 | 2088 | cytochrome P450, subfamily IIE (ethanol- | 22.58 | down | 0.00935 |
| J02888 at | 302888 | 2089 | NAD(P)H menadione oxidoreductase 2, dioxin- | 3.15 | down | 0.02385 |
| .102943 at | .102943 | 2090 | corticosteroid binding globulin | 18.98 | down | 0.00087 |
| .103242 s at | J03242 | 2092 | insulin-like growth factor 2 (somatomedin A) | 4.01 | down | 0.00042 |
| J03507 at | 103507 | 2095 | complement component 7 | 3.77 | down | 0.00184 |
| J03764 at | J03764 | 2097 | plasminogen activator inhibitor, type I | 5.6 | down | 0.02196 |
| l | | | protein phosphatase 2 (formerly 2A), catalytic | | | |
| J03805 s at | 303805 | 2098 | subunit, beta isoform | 3.87 | down | 0.0116 |
| J03810 at | J03810 | 2099 | solute carrier family 2 (facilitated glucose | 21.99 | down | 0.00004 |
| J03910 ma1 at | J03910 | 2101 | EST | 12.42 | down | 0.01167 |
| | | | methylenetetrahydrofolate dehydrogenase | | | |
| | | • | (NADP+ dependent), methenyltetrahydrofolate | | | |
| J04031 at | J04031 | 2103 | cyclohydrolase, formyltetrahydrofolate synthetase | 3.4 | down | 0.00786 |
| J04056_at | J04056 | 2104 | carbonyl reductase 1 | 5.19 | down | 0.00001 |
| J04080_at | J04080 | 2105 | complement component 1, s subcomponent | 5.48 | down | 0.0239 |
| J04093_s_at | J04093 | 2106 | UDP glycosyltransferase 1 | 18.92 | down | 0 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| at J04093 2106 UDP glycosyltransferase 1 at J0449 2110 oxidase), polypeptide 3 at J04615 2112 SNRPN upstream reading frame | Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|--|-------------|---------|--------|--|-------------|-----------|---------|
| 2110 oxidase), polypeptide 3 J04449 2110 oxidase), polypeptide 3 J04615 2112 SNRPN upstream reading frame syndecan 2 (heparan sulfate proteoglycan 1, cell J04621 2113 surface-associated, fibroglycan) J04621 2114 oxidase), polypeptide 5 J05037 2116 serine dehydratase J0518 2127 carboxypeptidase N, polypeptide B7 K02100 2125 component, Christmas disease, hemophilia B) K0216 2126 component, Christmas disease, hemophilia B) K0216 2127 component, Christmas disease, hemophilia B) K02176 2128 component Christmas disease, hemophilia B) K03192 2127 inducible), polypeptide 6 L00190 2130 antithrombin III L00352 2131 inducible), polypeptide 6 L004761 2138 oxidenmily IVA, polypeptide 11 L05779 2140 density ipoprotein receptor (familial L00972 2133 cystathionine-beta-synthase L07779 2140 epoxide hydrolase 2, cytoplasmic J140 2130 phosphoenolpyruvate carboxykinase 1 (soluble) L05779 2140 epoxide hydrolase 2, cytoplasmic J140 2140 epoxide hydrolase 2, cytoplasmic J141 2139 phosphoenolpyruvate carboxykinase 1 (soluble) L0777 2145 Coenzyme A dehydrogenase L07775 2147 carboxylesterase 1 (monocyte/macrophage | J04093 s at | J04093 | 2106 | UDP glycosyltransferase 1 | 18.92 | down | 0 |
| J04499 2110 oxidase), polypeptide 3 J04615 2112 SNRPN upstream reading frame syndecan 2 (heparan sulfate proteoglycan 1, cell J04621 2113 surface-associated, fibroglycan) cytochrome P450, subfamily IIIA (niphedipine J05037 2116 serine dahydratase J05158 2117 carboxypeptidase N, polypeptide 2, 83kD J05428 2120 UDP glycosyftransferase 2 family, polypeptide B7 K02100 2123 ornithine carbamoyltransferase K02215 2124 angiotensinogen coagulation factor IX (plasma thromboplastic K03192 2125 component, Christmas disease, hemophilia B) K03192 2127 inducible), polypeptide 6 L00190 2130 antithrombin III L00352 L00972 2131 inducible), polypeptide 6 L00972 2133 cystathionine-beta-synthase L0779 2140 apoxide hydrolase 2, cytoplasmic L0779 2140 apoxide hydrolase 2, cytoplasmic anyl-Coenzyme A, hydratase/3-hydroxyacyf L07077 2145 coenzyme A thydratase/3-hydroxyacyf L07775 2147 carboxylesterase 1 (monocyte/macrophage | l | | | cytochrome P450, subfamily IIIA (niphedipine | | | |
| J04615 2112 SNRPN upstream reading frame syndecan 2 (heparan sulfate proteoglycan 1, cell 3urface-associated, fibroglycan) cytochrome P450, subfamily IIIA (niphediphe J04813 2114 oxidase), polypeptide 5 J05037 2116 serine dehydratase Upby peptide 5 J17 carboxypeptidase N, polypeptide 2, 83kD J05428 2120 UDP glycosyltransferase 2 family, polypeptide B7 (2210 2123 ornithine carbamoyltransferase (J05037 2124 angiotensinogen coagulation factor IX (plasma thromboplastic cytochrome P450, subfamily IIA (phenobarbital-Inducible), polypeptide 6 cytophasmic 2133 phosphoenolpyruvate carboxykinase 1 (soluble) anoyl-Coenzyme A, hydratase/3-hydroxyacyf (coenzyme A, hydratase/3-hydroxyacyf (coenzyme A, hydratase/3-hydroxyacyf (coenzyme A, hydratase/3-hydroxyacyf (monocyfe/macrophage | J04449 at | J04449 | 2110 | oxidase), polypeptide 3 | 5.25 | down | 0.01583 |
| syndecan 2 (heparan sulfate proteoglycan 1, cell J04621 2113 surface-associated, fibroglycan) cytochrome P450, subfamily IIIA (niphedipine J04813 2114 oxidase), polypeptide 5 J05037 2116 serine dehydratase J05128 2120 UDP glycosytransferase 2 family, polypeptide B7 K02100 2123 omithine carbamoyltransferase 2 family, polypeptide B7 cagulation factor IX (plasma thromboplastic coagulation factor IX (plasma thromborbitial inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital L00352 2131 low density lipoprotein receptor (familial L00972 2133 cystathionine-beta-synthase L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05779 2144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) epoxide hydrotoxymethylglutaryl-Coenzyme A lyase (hydroxymethylglutaryl-Coenzyme A lyase (hydroxymethylglutaryl-Coenzyme A lyase (hydroxymethylglutaryl-Coenzyme A cenyme A dehydrogensse L07755 2147 carboxylesterase 1 (monocyfe/macrophage | J04615_at | J04615 | 2112 | SNRPN upstream reading frame | 3.14 | down | 0.02928 |
| J04621 2113 surface-associated, fibroglycan) cytochrome P450, subfamily IIIA (niphedipine J04813 2114 oxidase), polypeptide 5 J0558 2117 carboxypeptidase N, polypeptide 2, 83kD J05428 2120 UDP glycosyltransferase 2 family, polypeptide B7 K02100 2123 ornithine carbamoyltransferase K02215 2124 angiotensinogen Coagulation factor IX (plasma thromboplastic Coagulation factor IX (plasma thrombophilia B) Cytochrome P450, subfamily IIA (phenobarbital- inducible), polypeptide 6 Cytochrome P450, subfamily IIA (phenobarbital- Co0352 2127 inducible), polypeptide 6 Cytochrome P450, subfamily IIA (phenobarbital- L00352 2131 low density lipoprotein receptor (familial L00972 2133 cystathionine-beta-synthase L04751 2138 cystathionine-beta-synthase L05779 2143 phosphoenolpyruvate carboxykinase 1 (soluble) L05779 2144 jase (hydroxymethylalutaricaciduria) enoyl-Coenzyme A hydratase/3-hydroxyacyl L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | I | | - | syndecan 2 (heparan sulfate proteoglycan 1, cell | | | |
| cytochrome P450, subfamily IIIA (niphedipine oxidase), polypeptide 5 J05037 J05158 Z116 Serine dehydratase J05158 Z120 UDP glycosyltransferase 2 family, polypeptide B7 K02100 Z123 angiotensinogen Caguilation factor IX (plasma thromboplastic component, Christmas disease, hemophilia B) K02766 Z125 Component, Christmas disease, hemophilia B) K02766 Z126 Component, Christmas disease, hemophilia B) K03192 Z127 Inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 Cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 L00190 Z130 C131 L00352 Z131 L004751 Z138 Cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 L004751 Z139 Cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 Cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 C100372 Z131 Cystathionine-beta-synthase L07779 Z140 Phydroxymethyl-3-methylglutaryl-Coenzyme A L07033 Z144 Iyase (hydroxymethyl-3-methylglutaryl-Coenzyme A L07077 Z145 Coenzyme A dehydrogenase L07765 Z147 Carboxylesterase 1 (monocyte/macrophage | J04621_at | J04621 | 2113 | surface-associated, fibroglycan) | 3.38 | down | 0.00275 |
| Jo4813 2114 oxidase), polypeptide 5 Jo5037 2116 serine dehydratase Jo5158 2117 carboxypeptidase N, polypeptide 2, 83kD Jo5428 2120 UDP glycosytransferase 2 family, polypeptide B7 K02100 2123 omittine carbamoyltransferase K02215 component, Christmas disease, hemophilia B) K02402 2125 component, Christmas disease, hemophilia B) K02766 2126 complement component 9 Cytochrome P450, subfamily IIA (phenobarbital- K03192 2127 inducible), polypeptide 6 Cytochrome P450, subfamily IIA (phenobarbital- K03192 2137 inducible), polypeptide 6 Cytochrome P450, subfamily IVA, polypeptide 11 C0352 2133 ontithrombin III C0352 2133 cystathionine-beta-synthase L04751 2139 phosphoenolpyruvate carboxykinase 1 (soluble) L05779 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethyl-3-methylglutaricaciduria) enoyl-Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | | | | cytochrome P450, subfamily IIIA (niphedipine | | | |
| J05037 2116 serine dehydratase J05158 2117 carboxypeptidase N, polypeptide 2, 83kD J05428 2120 UDP glycosyltransferase 2 family, polypeptide B7 K02100 2123 ornithine carbamoyltransferase K02215 2124 angiotensinogen coagulation factor IX (plasma thromboplastic coagulation factor IX (plasma thromboplilia B) cytochrome P450, subfamily IIA (phenobarbital- coagulation ill coagon 2137 inducible), polypeptide 6 cytochrome P450, subfamily IVA, polypeptide 11 coagon 2131 ow density ipoprotein receptor (familial coagon 2133 ovtochrome P450, subfamily IVA, polypeptide 11 coagon 2133 cytochrome P450, subfamily IVA, polypeptide 11 coagon 2133 cytochrome P450, subfamily IVA, polypeptide 11 coagon 2134 phosphoenolopyruvate carboxykinase 1 (soluble) coagulation in the coagon coagulation in the c | J04813 s at | J04813 | 2114 | oxidase), polypeptide 5 | 29.6 | down | 0.0107 |
| J05158 2117 carboxypeptidase N, polypeptide 2, 83kD J05428 2120 UDP glycosyltransferase 2 family, polypeptide B7 K02100 2123 ornithine carbamoyltransferase K02215 2124 angiotensinogen Coagulation factor IX (plasma thromboplastic Coagulation factor IX (plasma IX (plasma thromboplastic) factor IX (plasma thromboplastic) factor IX (plasma thromboplastic) factor IX (plasma thromboplasma thromboplastic) factor IX (plasma thromboplasma thromboplasma thromboplasma thromboplasma thromboplasma thromboplasma thromboplasm | J05037 at | J05037 | 2116 | serine dehydratase | 16.24 | down | 0.00015 |
| 105428 2120 UDP glycosyltransferase 2 family, polypeptide B7 K02100 2123 ornithine carbamoyltransferase K02215 2124 angiotensinogen coagulation factor IX (plasma thromboplastic Coagulation factor IX (plasma factor) familial L00352 2131 low density lipoprotein receptor (familial L00352 2131 low density lipoprotein receptor (familial L00372 2133 cystathionine-beta-synthase L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethylglutaricaciduria) enoyl-Coenzyme A, hydratase/3-hydroxyacyl coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | J05158_at | J05158 | 2117 | carboxypeptidase N, polypeptide 2, 83kD | 8.52 | down | 0 |
| K02215 2124 angiotensinogen coagulation factor IX (plasma thromboplastic coagulation factor IX (plasma thromboplastic K02402 2125 component, Christmas disease, hemophilia B) K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital- cytochrome P450, subfamily IIA (phenobarbital- inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital- cytochrome P450, subfamily IIA (phenobarbital- inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital- inducible), polyp | J05428 at | J05428 | 2120 | UDP glycosyltransferase 2 family, polypeptide B7 | 16.14 | down | 0.00563 |
| K02215 2124 angiotensinogen coagulation factor IX (plasma thromboplastic K02402 2125 component, Christmas disease, hemophilia B) K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital- cytochrome P450, subfamily IIA (phenobarbital- inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital- at L00190 2127 inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital- at L00352 2127 inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital- at L00352 2131 low density lipoprotein receptor (familial L004751 2133 cytochrome P450, subfamily IVA, polypeptide 11 L05779 2140 sphosphoenolpyruvate carboxykinase 1 (soluble) t L05779 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethyl-3-methylglutaricaciduria) enoyl-Coenzyme A dehydrogenase t C07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | K02100_at | K02100 | 2123 | ornithine carbamoyltransferase | 10.24 | down | 0.0000 |
| coagulation factor IX (plasma thromboplastic K02402 2125 component, Christmas disease, hemophilia B) K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital- K03192 2127 inducible), polypeptide 6 L00190 2130 antithrombin III L00352 2131 low density lipoprotein receptor (familial L00972 2133 cystathionine-beta-synthase L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05779 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A, hydratase/3-hydroxyacyl t L07775 2147 carboxylesterase 1 (monocyte/macrophage | K02215 at | K02215 | 2124 | angiotensinogen | 16.51 | down | 0.00006 |
| K02402 2125 component, Christmas disease, hemophilia B) K02766 2126 complement component 9 Cytochrome P450, subfamily IIA (phenobarbital- inducible), polypeptide 6 Cytochrome P450, subfamily IIA (phenobarbital- inducible), polypeptide 6 L00190 2137 inducible), polypeptide 6 L00352 2131 low density lipoprotein receptor (familial L00372 2131 low density lipoprotein receptor (familial L004751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) L05779 2140 poxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A, hydratase/3-hydroxyacyl t L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | i | | | coagulation factor IX (plasma thromboplastic | | | |
| K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital- at K03192 2127 inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital- at L00190 2130 antithrombin III L00352 2131 low density lipoprotein receptor (familial L00972 2133 cystathionine-beta-synthase L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05779 cytochrome P450, subfamilial cyto | K02402 at | K02402 | 2125 | component, Christmas disease, hemophilia B) | 28.81 | down | 0.00001 |
| cytochrome P450, subfamily IIA (phenobarbital- K03192 2127 inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital- K03192 2127 inducible), polypeptide 6 L00190 2130 antithrombin III L00352 2131 low density lipoprotein receptor (familial L00972 2133 cystathionine-beta-synthase L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05744 2139 phosphoenolpyruvate carboxykinase 1 (soluble) L05779 cytochrome P450, subfamily IVA, polypeptide 11 L05779 cytochrome P450, subfamily IVA, polypeptide 11 L05779 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A, hydratase/3-hydroxyacyl t L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | K02766_at | K02766 | 2126 | complement component 9 | 21.24 | down | 0 |
| K03192 2127 inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital- cytochrome P450, subfamily IIA (phenobarbital- at L00190 2137 inducible), polypeptide 6 L00352 2131 low density lipoprotein receptor (familial L00372 2133 cystathionine-beta-synthase L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) L05779 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | i | | | cytochrome P450, subfamily IIA (phenobarbital- | | | |
| cytochrome P450, subfamily IIA (phenobarbital- at L00190 2127 inducible), polypeptide 6 L00190 2130 antithrombin III L00352 2131 low density lipoprotein receptor (familial L00972 2133 cystathionine-beta-synthase L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) L05779 2140 phosphoenolpyruvate carboxykinase 1 (soluble) L07783 2144 lyase (hydroxymethyl-3-methylglutaryl-Coenzyme A L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | K03192 f_at | K03192 | 2127 | inducible), polypeptide 6 | 69.92 | down | 0 |
| K03192 2127 inducible), polypeptide 6 L00190 2130 antithrombin III L00352 2131 low density lipoprotein receptor (familial L00972 2133 cystathionine-beta-synthase L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) L05779 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A hydratase/3-hydroxyacyl t L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | l I | | | cytochrome P450, subfamily IIA (phenobarbital- | | | |
| L00190 2130 antithrombin III L00352 2131 low density lipoprotein receptor (familial L00972 2133 cystathionine-beta-synthase L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) L05779 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A hydratase/3-hydroxyacyl L07077 2145 Coenzyme A dehydrogenase L077765 2147 carboxylesterase 1 (monocyte/macrophage | K03192 f at | K03192 | 2127 | inducible), polypeptide 6 | 50.16 | down | 0 |
| L00352 2131 low density lipoprotein receptor (familial L00972 2133 cystathionine-beta-synthase L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) copoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A dehydrogenase L07775 2145 Coenzyme A dehydrogenase to C07765 2147 carboxylesterase 1 (monocyte/macrophage | L00190 s at | L00190 | 2130 | antithrombin III | 42.41 | down | 0.00012 |
| L00972 2133 cystathionine-beta-synthase L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) L05779 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A, hydratase/3-hydroxyacyl L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | L00352 at | L00352 | 2131 | low density lipoprotein receptor (familial | 4.19 | down | 0.00352 |
| L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) L05779 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A, hydratase/3-hydroxyacyl L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | L00972_at | L00972 | 2133 | cystathionine-beta-synthase | 7.19 | down | 0.00008 |
| L05779 2140 epoxide hydrolase 2, cytoplasmic 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A, hydratase/3-hydroxyacyl L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | L04751 at | L04751 | 2138 | cytochrome P450, subfamily IVA, polypeptide 11 | 36.79 | down | 0.00004 |
| L05779 2140 epoxide hydrolase 2, cytoplasmic 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A, hydratase/3-hydroxyacyl L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | L05144 at | L'05144 | 2139 | phosphoenolpyruvate carboxykinase 1 (soluble) | 4.76 | down | 0.02289 |
| 3-hydroxymethyl-3-methylglutaryl-Coenzyme A L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A, hydratase/3-hydroxyacyl L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | L05779 at | L05779 | 2140 | epoxide hydrolase 2, cytoplasmic | 5.35 | down | 0.00006 |
| L07033 2144 lyase (hydroxymethylglutaricaciduria) enoyl-Coenzyme A, hydratase/3-hydroxyacyl L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | 1 | | | 3-hydroxymethyl-3-methylglutaryl-Coenzyme A | | | |
| enoyl-Coenzyme A, hydratase/3-hydroxyacyl L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | L07033_at | L07033 | 2144 | lyase (hydroxymethylglutaricaciduria) | 3.49 | down | 0 |
| L07077 2145 Coenzyme A dehydrogenase L07765 2147 carboxylesterase 1 (monocyte/macrophage | | | | enoyl-Coenzyme A, hydratase/3-hydroxyacyl | | | |
| L07765 2147 carboxylesterase 1 (monocyte/macrophage | L07077_at | L07077 | 2145 | Coenzyme A dehydrogenase | 4.82 | down | 0.00403 |
| | L07765_at | L07765 | 2147 | carboxylesterase 1 (monocyte/macrophage | 20.53 | down | 0.00025 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change Direction | Direction | Pvalue |
|-------------|---------|--------|---|-----------------------|-----------|---------|
| | | | glucan (1,4-alpha-), branching enzyme 1 | | | |
| L07956_at | L07956 | 2148 | (glycogen branching enzyme, Andersen disease, fatty-acid-Coenzyme A ligase, long-chain 1,fatty- | 5.6 | down | 0.00029 |
| L09229 s at | L09229 | 2150 | acid-Coenzyme A ligase, long-chain 2 | 18.34 | down | 0.00016 |
| L09708 at | L09708 | 2152 | complement component 2 | 3.92 | down | 0.00693 |
| L09717 at | L09717 | 2153 | lysosomal-associated membrane protein 2 | 4.06 | down | 0.00034 |
| L11005 at | L11005 | 2154 | aldehyde oxidase 1 | 16.3 | down | 0.00065 |
| L11244 s at | L11244 | 2155 | complement component 4-binding protein, beta | 43.33 | down | 0 |
| L11244 s at | L11244 | 2155 | complement component 4-binding protein, beta | 12.03 | down | 0.0001 |
| L11708 at | L11708 | 2158 | hydroxysteroid (17-beta) dehydrogenase 2 | 5.99 | down | 0.01516 |
| L11931 at | L11931 | 2159 | serine hydroxymethyltransferase 1 (soluble) | 7.27 | down | 0.00041 |
| L12760 s at | L12760 | 2162 | phosphoenolpyruvate carboxykinase 1 (soluble) | 12.75 | down | 0.00035 |
| L13278 at | L13278 | 2163 | crystallin, zeta (quinone reductase) | 5.83 | down | 0.0034 |
| L15702 at | L15702 | 2165 | B-factor, properdin | 3.7 | down | 0.04693 |
| ; | | | cytochrome P450, subfamily IIC (mephenytoin 4- | | | |
| L16883 s at | L16883 | 2166 | hydroxylase), polypeptide 9 | 84.71 | down | 0.00327 |
| L17128 at | L17128 | 2167 | gamma-glutamyl carboxylase | 4.02 | down | 0.00096 |
| ı | | | phosphodiesterase 4A, cAMP-specific (dunce | | | |
| L20965_at | L20965 | 2175 | (Drosophila)-homolog phosphodiesterase E2) solute carrier family 10 (sodium/bile acid | 3.02 | down | 0.01177 |
| 1- 64000 | 1 04000 | 2476 | cotrong family member 1 | 13.18 | מאטט | 0.00155 |
| 127548 at | 1 22548 | 2178 | collade two XVIII alpha 1 | 3.87 | down | 0.0299 |
| 1.25340_dt | 1 25878 | 2183 | enoxide hydrolase 1. microsomal (xenobiotic) | 26.84 | down | 0 |
| 125880 s at | L25880 | 2184 | epoxide hydrolase 1, microsomal (xenobiotic) | 58.7 | down | 0.00013 |
| L27050 at | L27050 | 2186 | apolipoprotein F | 10.26 | down | 0.00026 |
| L29008 at | L29008 | 2189 | sorbitol dehydrogenase | 3.51 | down | 0.00825 |
| L29433 at | L29433 | 2191 | coagulation factor X | 7.74 | down | 0.00244 |
| L32140 at | L32140 | . 2192 | afamin | 17.31 | down | 0.00003 |
| L32179 at | L32179 | 2193 | arylacetamide deacetylase (esterase) | 23.83 | down | 0 |
| 1 | | | bile acid Coenzyme A: amino acid N- | | | |
| L34081_at | L34081 | 2199 | acyltransferase (glycine N-choloyltransferase) | 11.96 | down | 0.00008 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|-----------|---------|
| | | | glutamate-cysteine ligase (gamma- | | | |
| L35546_at | L35546 | 2203 | glutamylcysteine synthetase), regulatory (30.8kD) | 14.18 | down | 0.00018 |
| | | , | giudinate-cystemia iigase (gamina- | ; | | |
| L35546_at | L35546 | 2203 | glutamylcysteine synthetase), regulatory (30.8kD) | 5.56 | down | 0.0005 |
| . L36033_at | L36033 | 2204 | stromal cell-derived factor 1 | 5.1 | down | 0.00603 |
| L38490_s_at | L38490 | 2207 | ADP-ribosylation factor 4-like | 3.13 | down | 0.01306 |
| | | | 5,10-methenyltetrahydrofolate synthetase (5- | | | |
| L38928 at | L38928 | 2209 | formyltetrahydrofolate cyclo-ligase) | 10.97 | down | 0.0267 |
| I | | | 5,10-methenyltetrahydrofolate synthetase (5- | | | |
| L38928_at | L38928 | 2209 | formyltetrahydrofolate cyclo-ligase) | 5.17 | down | 0.00726 |
| L40401 at | L40401 | 2211 | putative protein | 6.97 | down | 0.00079 |
| L40401 at | L40401 | 2211 | putative protein | 4.26 | down | 0.00194 |
| L41067 at | L41067 | 2213 | nuclear factor of activated T-cells, cytoplasmic 3 | 4.96 | down | 0.00473 |
| L47726_at | L47726 | 2219 | phenylalanine hydroxylase | 25.63 | down | 0.00019 |
| L48516_at | L48516 | 2220 | paraoxonase 3 | 22.21 | down | 0.00004 |
| L49169_at | L49169 | 2221 | FBJ murine osteosarcoma viral oncogene | 3.4 | down | 0.01193 |
| L76465_at | L76465 | 2224 | hydroxyprostaglandin dehydrogenase 15-(NAD) | 3.56 | down | 0.00688 |
| L76571_at | L76571 | 2226 | nuclear receptor subfamily 0, group B, member 2 | 4.44 | down | 0.00312 |
| L76687_at | L76687 | 2227 | growth factor receptor-bound protein 14 | 5.16 | down | 0.00199 |
| L76927_rna1_at | L76927 | 2228 | galactokinase 1 | 3.66 | down | 0.00999 |
| | | | solute carrier family 25 (mitochondrial carrier; | | | |
| L77567_s_at | L77567 | 2229 | citrate transporter), member 1 | 3.14 | down | 0.04095 |
| M10058_at | M10058 | 2230 | asialoglycoprotein receptor 1 | 23.96 | down | 0 |
| M10612_at | M10612 | 2232 | apolipoprotein C-II | 17.13 | down | 0.00746 |
| M10942_at | M10942 | 2233 | metallothionein 1E (functional) | 6.19 | down | 0.00428 |
| M10943_at | M10943 | 2234 | metallothionein 1F (functional) | 3.88 | down | 0 |
| M11025_s_at | M11025 | 2235 | asialoglycoprotein receptor 2 | 17.56 | down | 0.00003 |
| M11313_s_at | M11313 | 2236 | alpha-2-macroglobulin | 10.05 | down | 0.00014 |
| M11321_at | M11321 | 2237 | group-specific component (vitamin D binding | 16.52 | down | 0.01416 |
| M11437_cds1_at | M11437 | 2238 | kininogen | 18.38 | down | 0.00006 |
| M11437_cds2_at | M11437 | 2238 | kininogen | 16.19 | down | 0.02277 |
| 1 | | | | | | |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| 2 | Genbank | Sed ID | | rold Change | חופכווסוו | Lyaide |
|----------------|---------|--------|--|-------------|-----------|---------|
| M11567 ma1 at | M11567 | 2239 | angiogenin, ribonuclease, RNase A family, 5 | 32.25 | down | 0.0001 |
| M12174 at | M12174 | 2242 | ras homolog gene family, member B | 5.44 | down | 0.0088 |
| • | | | alcohol dehydrogenase 1 (class I), alpha | | | |
| | | | polypeptide, alcohol dehydrogenase 2 (class I), | | | |
| M12272 s at | M12272 | 2243 | beta polypeptide, alcohol dehydrogenase 3 (class | 32.42 | down | 0.0034 |
| M12529 at | M12529 | 2244 | apolipoprotein E | 3.05 | down | 0.03776 |
| M12625 at | M12625 | 2245 | lecithin-cholesterol acyltransferase | 4.55 | down | 0.01584 |
| M12712 s at | M12712 | 2246 | protein C (inactivator of coagulation factors Va | 7.37 | down | 0.01866 |
| 1 | | | alcohol dehydrogenase 1 (class I), alpha | | | |
| | | | polypeptide, alcohol dehydrogenase 2 (class I), | | | |
| M12963 s at | M12963 | 2248 | beta polypeptide, alcohol dehydrogenase 3 (class | 48.95 | down | 0.00104 |
| M13143 at | M13143 | 2249 | kallikrein B plasma, (Fletcher factor) 1 | 10.39 | down | 0.00019 |
| M13149 at | M13149 | 2250 | histidine-rich glycoprotein | 18.65 | down | 0.02974 |
| | M13232 | 2251 | coagulation factor VII (serum prothrombin | 5.9 | down | 0.00014 |
| M13690 s at | M13690 | 2252 | complement component 1 inhibitor (angioedema, | 6.07 | down | 0.00045 |
| M13699 at | M13699 | 2253 | ceruloplasmin (ferroxidase) | 15.85 | down | 0.00012 |
| M13829 s at | M13829 | 2254 | v-raf murine sarcoma 3611 viral oncogene | 6.52 | down | |
| M14058 at | M14058 | 2256 | complement component 1, r subcomponent | 99.9 | down | 0.00229 |
| M14091 at | M14091 | 2257 | thyroxin-binding globulin | 10.66 | down | 0.00024 |
| M14218 at | M14218 | 2259 | argininosuccinate lyase | 9.03 | down | 0.00078 |
| M14338 at | M14338 | 2260 | protein S (alpha) | 12.33 | down | 0 |
| l | | | phosphorylase, glycogen; liver (Hers disease, | | | |
| M14636 at | M14636 | 2262 | glycogen storage disease type VI) | 3.45 | down | 0.00133 |
| M15465 s at | M15465 | 2266 | pyruvate kinase, liver and RBC | 6.1 | down | 0.00069 |
| M15517 cds5 at | M15517 | 2267 | EST | 22.76 | down | 0.03365 |
| M15656 at | M15656 | 2268 | aldolase B, fructose-bisphosphate | 99.96 | down | 0 |
| M16447 at | M16447 | 2270 | quinoid dihydropteridine reductase | 6.57 | down | 0.00015 |
| M16474 s at | M16474 | 2271 | butyrylcholinesterase | 5.82 | down | 0.00113 |
| M16594 at | M16594 | 2272 | glutathione S-transferase A2 | 73.21 | down | |
| M16750 s at | M16750 | 2273 | pim-1 oncogene | 3.08 | down | 0.01811 |
| MARORA | M46064 | 2274 | alpha-2-HS-clycoprotein | 21.45 | מאיטף | 0.01175 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|-------------|---------|--------|---|-------------|-----------|---------|
| M16967_at | M16967 | 2275 | coagulation factor V (proaccelerin, labile factor) | 5.56 | down | 0.00047 |
| M16973_at | M16973 | 2276 | complement component 8, beta polypeptide | 22.75 | down | 0.00001 |
| M16974_s_at | M16974 | 2277 | complement component 8, alpha polypeptide | 49.47 | down | 0.00046 |
| M17262_at | M17262 | 2278 | coagulation factor II (thrombin) | 44.3 | down | 0.00345 |
| M17262_at | M17262 | 2278 | coagulation factor II (thrombin) | 14.24 | down | 0.00028 |
| M17466_at | M17466 | 2279 | coagulation factor XII (Hageman factor) | 9.76 | down | 0.00285 |
| i | | | dystrophin (muscular dystrophy, Duchenne and Becker types), includes DXS142, DXS164, | | | |
| M18533_at | M18533 | 2284 | DXS206, DXS230, DXS239, DXS268, DXS269, | 3.45 | down | 0.00313 |
| M19828_s_at | M19828 | 2287 | apolipoprotein B (including Ag(x) antigen) | 29.37 | down | 0.00137 |
| M20218 at | M20218 | 2288 | coagulation factor XI (plasma thromboplastin | 6.4 | down | 0.00004 |
| M20786_at | M20786 | 2290 | alpha-2-plasmin inhibitor | 16.95 | down | 0.00709 |
| M20867_s_at | M20867 | 2291 | glutamate dehydrogenase 1 | 17.73 | down | 0.00002 |
| M20902_at | M20902 | 2292 | apolipoprotein C-l | 3.14 | down | 0.0389 |
| M21642_at | M21642 | 2294 | antithrombin III | 15.82 | down | 0.01027 |
| M21642_s_at | M21642 | 2294 | antithrombin III | 15.23 | down | 0.02088 |
| M22976_at | M22976 | 2297 | cytochrome b-5 | 7.39 | down | 0.02431 |
| M2316:1_at | M23161 | 2298 | EST | 3.44 | down | 0.00733 |
| M23234_s_at | M23234 | 2299 | ATP-binding cassette, sub-family B (MDR/TAP), | 10.05 | down | 0 |
| | | | androgen receptor (dihydrotestosterone receptor; | | | |
| | | | testicular feminization; spinal and bulbar | | | |
| M23263_at | M23263 | 2300 | muscular atrophy; Kennedy disease) | 4.6 | down | 0.00005 |
| | | | androgen receptor (dihydrotestosterone receptor; testicular feminization; spinal and bulbar | | | • |
| M23263 at | M23263 | 2300 | muscular atrophy; Kennedy disease) | 3.35 | down | 0.02551 |
| l | | | intercellular adhesion molecule 1 (CD54), human | | | |
| M24283_at | M24283 | 2303 | rhinovirus receptor | 3.19 | down | 0.04985 |
| M25079 s at | M25079 | 2305 | hemoglobin, beta | 9.15 | down | 0.01399 |
| M25280 at | M25280 | 2306 | selectin L (lymphocyte adhesion molecule 1) | 3.39 | down | 0.004 |
| M26393_s_at | M26393 | 2309 | | 16.27 | down | 0.00007 |
| M27492_at | M27492 | 2312 | | 4.62 | down | 0.0082 |
| | | | | | | |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| ## M29194 2315 lipase, hepatic ## M29873 2318 cytochrome P450, subfamily IIB (phenobarbital- ## M29874 2329 cytochrome P450, subfamily IIB (phenobarbital- ## M29974 2320 cytochrome P450, subfamily IIB (phenobarbital- ## M30185 2321 cholesteryl ester transfer protein, plasma ## M30267 2322 cholesteryl ester transfer protein, plasma ## M30267 2323 propionyl Coenzyme A carboxylase, beta ## M3169 2322 propionyl Coenzyme A carboxylase, beta ## M3169 2325 propionyl Coenzyme A carboxylase, beta ## M31607 2330 propionyl Coenzyme A carboxylase, beta ## M31607 2331 compound-inducible), polypeptide 2 ## M31994 2332 aldehyde dehydrogenase 1, soluble ## M33317 2338 inducible), polypeptide 7 ## M33518 2339 inducible), polypeptide 7 ## M35590 2345 small inducible cytokine A4 ## M35590 2345 small inducible cytokine A4 ## M35590 2345 small inducible cytokine A4 ## M35690 2348 (aspartate aminotransferase 1) ## M55691 2352 protein Z, vitamin K-dependent plasma ## M5561 2359 protein Z, vitamin K-dependent plasma ## M5561 2359 protein Z, vitamin K-dependent plasma ## M5561 2350 protein Z, vitamin K-dependent plasma ## M55786 2360 protein Z, vitamin K-dependent plasma ## M55786 2360 protein Z, vitamin K-dependent plasma ## M55781 2359 GROZ oncogene | Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue | |
|--|-------------|---------|--------|---|-------------|-----------|---------|--|
| M29873 2318 cytochrome P450, subfamily IIB (phenobarbital-N29874 5318 cytochrome P450, subfamily IIB (phenobarbital-N2320 0-6-methylguanine-DNA methyltransferase 3.97 M30185 2321 cholesteryl easter transfer protein, plasma 3.39 M30287 2322 vascular cell adhesion molecule 1 3.41 M30289 2323 indogen (enactin) 3.4 M31697 2330 x-box binding protein 1 3.4 M31697 2331 compound-inducible), polypeptide 2 4.38 M31667 2332 aldehyde dehydrogenase 1, soluble 4.38 M31994 2332 aldehyde dehydrogenase 1, soluble 4.38 M33317 2338 inducible), polypeptide 7 4.38 M33318 2332 aldehyde dehydrogenase 1, soluble 11.24 M35400 2344 insulin-like growth factor binding protein 2 (36kD) 3.65 M35590 2344 insulin-like growth factor binding protein 3 24.73 M35590 2345 small inducible cydokine A4 4.74 M35590 2345 small inducible | 29194_at | M29194 | 2315 | lipase, hepatic | 12.18 | down | 0.00012 | |
| M29874 2319 cytochrome P450, subfamily IIB (phenobarbital- 18.44 M29871 2320 O-6-methylguanine-DNA methyltransferase 3.97 M30185 2321 cholesteryl ester transfer protein, plasma 3.39 M30269 2322 vacocular cell adhesion molecule 1 3.4 M31699 2325 propionyl Coenzyme A carboxylase, beta 4.65 M31697 2330 x-box binding protein 1 3.4 M31697 2331 compound-inducible), polypeptide 2 4.38 M31697 2332 aldocible), polypeptide 2 4.38 M31994 2332 inducible), polypeptide 2 4.38 M31994 2332 inducible), polypeptide 7 4.38 M33317 2338 inducible), polypeptide 6 4.38 M3477 2344 plasminogan 3.21 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.38 M3550 | 9873_s_at | M29873 | 2318 | cytochrome P450, subfamily IIB (phenobarbital- | 56.71 | down | 0.0054 | |
| M29971 2320 O-6-methylguanine-DNA methyltransferase 3.97 M30185 2321 cholesteryl ester transfer protein, plasma 3.83 M30185 2321 cholesteryl ester transfer protein, plasma 3.39 M30257 2322 vascular cell adhesion molecule 1 3.41 M31699 2325 propionyl Coenzyme A carboxylase, beta 4.65 M31627 2330 X-box binding protein 1 3.41 Cytochrome P450, subfamily Il (aromatic cytochrome P450, subfamily IlA (phenobarbital cytochrome P450, subfamily IIA (| 9874 s at | M29874 | 2319 | cytochrome P450, subfamily IIB (phenobarbital- | 18.44 | down | 0.00081 | |
| M30185 2321 cholesteryl ester transfer protein, plasma 3.83 M30269 2322 cholesteryl ester transfer protein, plasma 3.39 M30269 2322 vascular cell adhesion molecule 1 3.11 M3169 2325 propionyl Coenzyme A carboxylase, beta 4.65 M3167 2330 X-box binding protein 1 4.65 M3167 2331 compound-inducible), polypeptide 2 4.38 M31894 2332 aldehyde dehydrogenase 1, soluble 11.24 cytochrome P450, subfamily IIA (phenobarbital-dytochrome P450, subfamily IIA (phenobarbital-dytochrome P450, subfamily IIA (phenobarbital-lytochrome P45 | 29971_at | M29971 | 2320 | O-6-methylguanine-DNA methyltransferase | 3.97 | down | 0.00424 | |
| M30185 2321 cholesteryl ester transfer protein, plasma 3.39 M30257 2322 vascular cell adhesion molecule 1 3.11 M30269 2322 protein of cenactin) 3.4 M3169 2325 propionyl Coenzyme A carboxylase, beta 4.65 M3167 2330 X-box binding protein 1 6.37 cytochrome P450, subfamily I (aromatic 6.37 M31994 2332 aldehyde dehydrogenase 1, soluble 11.24 cytochrome P450, subfamily IIA (phenobarbital-dytochrome P450 | 30185_at | M30185 | 2321 | cholesteryl ester transfer protein, plasma | 3.83 | down | 0.0013 | |
| M30257 2322 vascular cell adhesion molecule 1 3.11 M30269 2323 nidogen (enactin) 3.4 M3169 2325 propionyl Coenzyme A carboxylase, beta 4.65 M31627 2330 X-box binding protein 1 4.65 M31667 2331 compound-inducible), polypeptide 2 4.38 M31994 2332 aldehyde dehydrogenase 1, soluble 4.38 Cytochrome P450, subfamily IIA (phenobarbital-cytochrome P450, subfamily IIA (phenobarbital-cytochrome P450, subfamily IIA (phenobarbital-state) 32.63 M33318 2339 inducible), polypeptide 6 3247 M35410 2341 plasminogen 24.73 M35590 2345 inducible, polypeptide 6 24.73 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.62 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.74 M35590 2346 insulin-like growth factor binding protein 3 4.35 M55150 2352 tumarylacetoacetate 3.93 </td <td>30185_at</td> <td>M30185</td> <td>2321</td> <td>cholesteryl ester transfer protein, plasma</td> <td>3.39</td> <td>down</td> <td>0.00089</td> <td></td> | 30185_at | M30185 | 2321 | cholesteryl ester transfer protein, plasma | 3.39 | down | 0.00089 | |
| M30269 2323 nidogen (enactin) 3.4 M3169 2325 propionyl Coenzyme A carboxylase, beta 4.65 M31627 235 X-box binding protein 1 4.65 M31667 233 compound-inducible), polypeptide 2 4.38 M31994 2332 aldehyde dehydrogenase 1, soluble 11.24 cytochrome P450, subfamily IIA (phenobarbital-cytochrome P450, subfamily IIA (phenobarbital-dytochrome P450, subfamily IIA (phenobarbital-lytochrome P450, subfamily IIA (phenobarbital-dytochrome P450, subfa | 0257 s at | M30257 | 2322 | vascular cell adhesion molecule 1 | 3.11 | down | 0.00064 | |
| M31169 2325 propionyl Coenzyme A carboxylase, beta 4.65 M31627 2330 X-box binding protein 1 6.97 Cytochrome P450, subfamily I (aromatic 4.38 M31667 2331 compound-inducible), polypeptide 2 4.38 M31994 2332 aldehyde dehydrogenase 1, soluble 11.24 Cytochrome P450, subfamily IIA (phenobarbital-oytochrome P450, subfamily IIA (phenobarbit | 30269 at | M30269 | 2323 | nidogen (enactin) | 3.4 | down | 0.00026 | |
| M31627 2330 X-box binding protein 1 6.97 cyfochrome P450, subfamily I (aromatic 4.38 M31994 2332 aldehyde dehydrogenase 1, soluble 11.24 M31994 2332 aldehyde dehydrogenase 1, soluble 11.24 M33317 2338 inducible), polypeptide 7 32.63 M33318 2339 inducible), polypeptide 6 32.13 M34276 2341 plasminogen 24.73 M34276 2343 insulin-like growth factor binding protein 2 (36kD) 6.45 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.74 M35570 2345 small inducible cytokine A4 4.62 M35570 2345 small inducible cytokine A4 4.62 M35570 2345 small inducible cytokine A4 4.62 M35670 2348 (asparatate aminotransferase 1) 4.37 M55150 2352 fumarylacetoacetate 5.96 M55513 2354 subfamily, member 5 | 1169 s at | M31169 | 2325 | propionyl Coenzyme A carboxylase, beta | 4.65 | down | 0.00467 | |
| Cytochrome P450, subfamily I (aromatic cytochrome P450, subfamily I (aromatic cytochrome P450, subfamily IIA (phenobarbital-cytochrome P450, subfamily IIA (phenobarbital-cytorhrome P450, subfamily IIA (phen | 31627_at | M31627 | 2330 | X-box binding protein 1 | 6.97 | down | 0.00052 | |
| M31667 2331 compound-inducible), polypeptide 2 4.38 M31994 2332 aldehyde dehydrogenase 1, soluble 11.24 cytochrome P450, subfamily IIA (phenobarbital-m33318 2338 inducible), polypeptide 7 32.63 cytochrome P450, subfamily IIA (phenobarbital-m33318 2339 inducible), polypeptide 6 3.21 M34276 2341 plasminogen m34276 24.73 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.74 M35670 2345 small inducible cytokine A4 4.74 M35878 2346 insulin-like growth factor binding protein 3 4.3 M35879 2346 insulin-like growth factor binding protein 3 4.3 M35878 2346 insulin-like growth factor binding protein 3 4.3 M35150 2348 (aspartate aminotransferase 1) 8.7 M55150 2352 fumarylacetoacetate 4.81 M55671 2354 subfamily, member 5 4.74 M55731 2359 | | | | cytochrome P450, subfamily I (aromatic | | | | |
| M31994 2332 aldehyde dehydrogenase 1, soluble 11.24 cytochrome P450, subfamily IIA (phenobarbital-dytochrome P450, subfamily IIA (phenobarbital-subderhole), polypeptide 6 32.63 M33318 2339 inducible), polypeptide 6 3.21 M34276 2341 plasminogen 24.73 M35590 2345 small inducible cytokine A4 24.74 M35590 2345 small inducible cytokine A4 4.74 M35690 2345 small inducible cytokine A4 4.74 M35878 2346 insulin-like growth factor binding protein 3 4.3 M35879 2348 insulin-like growth factor binding protein 3 4.3 M35879 2348 insulin-like growth factor binding protein 3 4.3 M35879 2348 insulin-like growth factor binding protein 3 4.3 M35879 2348 (aspartate aminotransferase 1) 8.7 M55150 2352 fumarylacetoacetate 3.93 potassium voltage-gated channel, shaker-related 4.81 M55671 2354 subfamilly, member 5 4.74 M57731 2359 protein Z, vitamin K-dependent plasma 4.74 M58286 2360 tumor necrosis factor receptor superfamily. | 31667 f at | M31667 | 2331 | compound-inducible), polypeptide 2 | 4.38 | down | 0.00078 | |
| A33317 2338 inducible), polypeptide 7 Cytochrome P450, subfamily IIA (phenobarbital- A33318 2339 inducible), polypeptide 6 A34276 2341 plasminogen A35590 2345 small inducible cytokine A4 A35590 2345 subfamily, member 5 A355 fumarylacetoacetate A355 protein Z, vitamin K-dependent plasma A3740 2355 protein Z, vitamin K-dependent plasma A37731 2359 GRO2 oncogene A375 A585 2360 tumor necrosis factor receptor superfamily. B15 | 131994 at | M31994 | 2332 | aldehyde dehydrogenase 1, soluble | 11.24 | down | 0.01192 | |
| M33317 2338 inducible), polypeptide 7 32.63 cytochrome P450, subfamily IIA (phenobarbital-M33318 2339 inducible), polypeptide 6 3.21 M34276 2341 plasminogen 24.73 M35410 2344 insulin-like growth factor binding protein 2 (36kD) 6.45 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.74 M35700 2346 insulin-like growth factor binding protein 3 4.3 glutamic-oxaloacetic transaminase 1, soluble 8.7 M37400 2348 (aspartate aminotransferase 1) 8.7 M55150 2352 fumarylacetoacetate 3.93 potassium voltage-gated channel, shaker-related 4.81 M5513 2354 subfamily, member 5 4.74 M55731 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 ctocogene 4.74 M58286 2360 tumor necrosis factor receptor superfamily. 8.15 | Ĭ | | | cytochrome P450, subfamily IIA (phenobarbital- | | | | |
| cytochrome P450, subfamily IIA (phenobarbital- M33318 2339 inducible), polypeptide 6 M354276 2341 plasminogen M355410 2344 insulin-like growth factor binding protein 2 (36kD) 6.45 M35590 2345 small inducible cytokine A4 M35590 2345 small inducible cytokine A4 M35590 2346 insulin-like growth factor binding protein 3 glutamic-oxaloacetic transaminase 1, soluble M37400 2348 (asparfate aminotransferase 1) M55150 2352 fumarylacetoacetate potassium voltage-gated channel, shaker-related M55513 2354 subfamily, member 5 M55731 2359 GRO2 oncogene M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | 33317 f at | M33317 | 2338 | inducible), polypeptide 7 | 32.63 | down | 0 | |
| M33318 2339 inducible), polypeptide 6 3.21 M34276 2341 plasminogen 24.73 M35410 2344 insulin-like growth factor binding protein 2 (36kD) 6.45 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.74 M35578 2346 insulin-like growth factor binding protein 3 4.3 M37400 2348 (aspartate aminotransferase 1) 8.7 M55150 2352 fumarylacetoacetate 8.7 M55150 2352 fumarylacetoacetate 3.93 M55513 2354 subfamily, member 5 4.81 M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 protein Z, vitamin K-dependent plasma 4.74 M58286 2360 tumor necrosis factor receptor superfamily. 8.15 | l I | | | cytochrome P450, subfamily IIA (phenobarbital- | | | , | |
| M34276 2341 plasminogen 24.73 M35410 2344 insulin-like growth factor binding protein 2 (36kD) 6.45 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.74 M35590 2345 small inducible cytokine A4 4.74 M35878 2346 insulin-like growth factor binding protein 3 4.3 glutamic-oxaloacetic transaminase 1, soluble 4.3 M37400 2348 (aspartate aminotransferase 1) 8.7 M55150 2352 fumarylacetoacetate 3.93 potassium voltage-gated channel, shaker-related 4.81 M55513 2354 subfamily, member 5 4.81 M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 protein Z, vitamin K-dependent plasma 4.74 M58286 2360 tumor necrosis factor receptor superfamily. 8.15 | 3318 r at | M33318 | 2339 | inducible), polypeptide 6 | 3.21 | down | 0.01621 | |
| M35410 2344 insulin-like growth factor binding protein 2 (36kD) 6.45 M35590 2345 small inducible cytokine A4 5.96 M35590 2345 small inducible cytokine A4 4.74 M35578 2346 insulin-like growth factor binding protein 3 4.62 M37400 2348 (aspartate aminotransferase 1) 8.7 M55150 2352 fumarylacetoacetate 8.7 M55513 2354 subfamily, member 5 4.81 M55573 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 chroein Z, vitamin K-dependent plasma 4.74 M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | 34276 at | M34276 | 2341 | plasminogen | 24.73 | down | 0.00031 | |
| M35590 2345 small inducible cytokine A4 5.96 M35590 2345 small inducible cytokine A4 4.74 M35878 2346 insulin-like growth factor binding protein 3 4.3 M35878 2346 insulin-like growth factor binding protein 3 4.3 M35878 2348 (aspartate aminotransferase 1) 8.7 M35150 2352 fumarylacetoacetate 8.7 M55150 2352 fumarylacetoacetate 3.93 M55513 2354 subfamily, member 5 4.81 M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 protein Z, vitamin K-dependent plasma 4.74 M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | 5410 s at | M35410 | 2344 | insulin-like growth factor binding protein 2 (36kD) | 6.45 | down | 0.04517 | |
| M35590 2345 small inducible cytokine A4 4.74 M35878 2346 insulin-like growth factor binding protein 3 4.62 M35878 2346 insulin-like growth factor binding protein 3 4.3 M37400 2348 (aspartate aminotransferase 1) 8.7 M55150 2352 fumarylacetoacetate 3.93 potassium voltage-gated channel, shaker-related 4.81 M55513 2354 subfamily, member 5 4.81 M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 GRO2 oncogene 4.74 M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | IP1-B at | M35590 | 2345 | small inducible cytokine A4 | 5.96 | down | 0.00604 | |
| M35590 2345 small inducible cytokine A4 4.62 M35878 2346 insulin-like growth factor binding protein 3 4.3 glutamic-oxaloacetic transaminase 1, soluble 4.3 M37400 2348 (aspartate aminotransferase 1) 8.7 M55150 2352 fumarylacetoacetate 3.93 potassium voltage-gated channel, shaker-related 4.81 M55513 2354 subfamily, member 5 4.81 M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 GRO2 oncogene 4.74 M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | IP1-B at | M35590 | 2345 | small inducible cytokine A4 | 4.74 | down | 0.01225 | |
| M35878 2346 insulin-like growth factor binding protein 3 4.3 glutamic-oxaloacetic transaminase 1, soluble M37400 2348 (aspartate aminotransferase 1) 8.7 M55150 2352 furnarylacetoacetate potassium voltage-gated channel, shaker-related M55513 2354 subfamily, member 5 M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 GRO2 oncogene M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | IIP1-B_at | M35590 | 2345 | small inducible cytokine A4 | 4.62 | down | 0.01268 | |
| glutamic-oxaloacetic transaminase 1, soluble M37400 2348 (aspartate aminotransferase 1) M55150 2352 fumarylacetoacetate potassium voltage-gated channel, shaker-related M55513 2354 subfamily, member 5 M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 GRO2 oncogene M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | 35878_at | M35878 | 2346 | insulin-like growth factor binding protein 3 | 4.3 | down | 0.0027 | |
| M37400 2348 (aspartate aminotransferase 1) 8.7 M55150 2352 fumarylacetoacetate potassium voltage-gated channel, shaker-related M55513 2354 subfamily, member 5 M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 GRO2 oncogene M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | | | | glutamic-oxaloacetic transaminase 1, soluble | | | | |
| M55150 2352 fumarylacetoacetate 3.93 potassium voltage-gated channel, shaker-related M55513 2354 subfamily, member 5 M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 GRO2 oncogene M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | 137400_at | M37400 | 2348 | (aspartate aminotransferase 1) | 8.7 | down | 0.0004 | |
| potassium voltage-gated channel, shaker-related M55513 2354 subfamily, member 5 M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 GRO2 oncogene M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | 155150 at | M55150 | 2352 | fumarylacetoacetate | 3.93 | down | 0.00213 | |
| M55513 2354 subfamily, member 5 M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 GRO2 oncogene 13.87 M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | l | | | potassium voltage-gated channel, shaker-related | | | | |
| M55671 2355 protein Z, vitamin K-dependent plasma 4.74 M57731 2359 GRO2 oncogene 13.87 M58286 2360 tumor necrosis factor receptor superfamily, 8.15 | 5513_s_at | M55513 | 2354 | subfamily, member 5 | 4.81 | down | 0.02141 | |
| at M58286 2360 tumor necrosis factor receptor superfamily. 8.15 | 55671 at | M55671 | 2355 | protein Z, vitamin K-dependent plasma | 4.74 | down | 0.00078 | |
| at M58286 2360 tumor necrosis factor receptor superfamily. 8.15 | | M57731 | 2359 | GRO2 oncogene | 13.87 | down | 0.0123 | |
| | M58286_s_at | M58286 | 2360 | tumor necrosis factor receptor superfamily, | 8.15 | down | 0.00037 | |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank Seq ID | <u>♀</u> | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------------|------------------------|--|-------------|-----------|---------|
| M58569 s at | M58569 2361 | 51 EST | | 16.15 | down | 0.00174 |
| M58600 rna1 at | M58600 . 2362 | 52 heparin cofactor II | factor II | 39.79 | down | 0.00034 |
| 1 | | tissue fact | tissue factor pathway inhibitor (lipoprotein- | • | | |
| M59499_at | M59499 2363 | | associated coagulation inhibitor) | 5.92 | down | 0.0005 |
| M59815 at | M59815 23 | 2364 compleme | complement component 4A | 8.02 | down | 0.00049 |
| | | sphingom | sphingomyelin phosphodiesterase 1, acid | | | |
| M59916 at | M59916 2365 | _ | ysosomal (acid sphingomyelinase) | 3.36 | down | 0.0038 |
| M60974 s at | M60974 2368 | _ | growth arrest and DNA-damage-inducible, alpha | 3.48 | down | 0.00209 |
| l I | | cytochrom | cytochrome P450, subfamily IIC (mephenytoin 4- | | | |
| M61853 at | M61853 2369 | _ | nydroxylase), polypeptide 18 | 7.82 | down | 0.00024 |
| M61854 s at | M61854 2370 | Ī | cytochrome P450, subfamily IIC (mephenytoin 4- | 3.3 | down | 0.04185 |
| I ! | | Ī | cytochrome P450, subfamily IIC (mephenytoin 4- | | | |
| M61855 at | M61855 2371 | | hydroxylase), polypeptide 9 | 38.82 | down | 0.00023 |
| M62403 s at | M62403 23 | 2373 insulin-like | insulin-like growth factor-binding protein 4 | 4.12 | down | 0.00226 |
| M62486_at | M62486 23 | 2374 compleme | complement component 4-binding protein, alpha | 22.08 | down | 0.00272 |
| ŀ | | glutathione | glutathione S-transferase M1, glutathione S- | | | |
| M63509 s at | M63509 23 | 2376 transferas | transferase M2 (muscle),glutathione S- | 7.06 | down | 0.03887 |
| M63967 at | M63967 23 | 2378 aldehyde o | aldehyde dehydrogenase 5 | 4.04 | down | 0.00058 |
| M64554_rna1_at | M64554 23 | 2380 coagulatio | coagulation factor XIII, B polypeptide | 5.87 | down | 0.00011 |
| i I | | glycine de | glycine dehydrogenase (decarboxylating; glycine | | | |
| M64590_at | M64590 23 | 2381 decarboxy | decarboxylase, glycine cleavage system protein | 6.41 | down | 0.00002 |
| M65131 rna1 at | M65131 23 | 2384 methylmal | methylmalonyl Coenzyme A mutase | 7.44 | down | 0.00004 |
| M65134 s at | , | 2385 compleme | complement component 5 | 12.01 | down | 0.00012 |
| M65292 s at | M65292 23 | 2386 H factor (c | H factor (complement)-like 1,H factor 1 | 7.56 | down | 0.01152 |
| M68516 rna1 at | M68516 23 | 2387 protein C i | protein C inhibitor (plasminogen activator inhibitor | 20.54 | down | 0 |
| M68840 at | | 2388 monoamir | monoamine oxidase A | 3.96 | down | 0.01396 |
| M68895 rna1_at | | 2390 alcohol de | alcohol dehydrogenase 6 (class V) | 4.25 | down | 0.00354 |
| M69177 at | | 2392 monoamir | monoamine oxidase B | 11.64 | down | 0.00001 |
| M72885 rna1 s at | | 2393 putative ly | putative lymphocyte G0/G1 switch gene | 6.5 | down | 0.03461 |
| M74587 rna1 s at | M74587 23 | 2394 insulin-like | insulin-like growth factor binding protein 1 | 11.42 | down | 0.00274 |
| M75106_at | M75106 23 | 2397 carboxype | carboxypeptidase B2 (plasma) | 40.63 | down | 0 |
| | | | | | | |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Sea ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|-----------|---------|
| M76665 at | M76665 | 2398 | hydroxysteroid (11-beta) dehydrogenase 1 | 19.22 | down | 0.00004 |
| M80482_at | M80482 | 2403 | paired basic amino acid cleaving system 4 | 4.26 | down | 0.00041 |
| M81182 s at | M81182 | 2404 | ATP-binding cassette, sub-family D (ALD), | 3.45 | down | 0.00499 |
| M81349 at | M81349 | 2405 | serum amyloid A4, constitutive | 76.15 | down | 0.00015 |
| M83216 s at | M83216 | 2407 | caldesmon 1 | 4.27 | down | 0.00037 |
| M83652 s at | M83652 | 2408 | properdin P factor, complement | 9 | down | 0.00002 |
| M83772 at | M83772 | 2409 | flavin containing monooxygenase 3 | 19.54 | down | 0 |
| M86826_at | M86826 | 2413 | insulin-like growth factor binding protein, acid | 3.75 | down | 0.01157 |
| M86873 s at | M86873 | 2414 | plasminogen,plasminogen-like | 17.54 | down | 0 |
| 1 | | | SWI/SNF related, matrix associated, actin | | | |
| M88163 at | M88163 | 2416 | dependent regulator of chromatin, subfamily a, | 3.37 | down | 0.00098 |
| M91432_at | M91432 | 2420 | acyl-Coenzyme A dehydrogenase, C-4 to C-12 | 6.74 | down | 0.00008 |
| M92843 s at | M92843 | 2421 | zinc finger protein homologous to Zfp-36 in | 3.02 | down | 0.04958 |
| M93143 at | M93143 | 2423 | plasminogen-like | 10.06 | down | 0.00098 |
| M93405 at | M93405 | 2424 | methylmalonate-semialdehyde dehydrogenase | 23.06 | down | 0 |
| M94065 s at | M94065 | 2425 | dihydroorotate dehydrogenase | 11.78 | down | 0.00034 |
| M94065 at | M94065 | 2425 | dihydroorotate dehydrogenase | 6.47 | down | 0.00013 |
| M95585 s at | M95585 | 2430 | hepatic leukemia factor | 4.2 | down | 0.00212 |
| M95767 at | M95767 | 2432 | chitobiase, di-N-acetyl- | 4.94 | down | 0.00004 |
| | | | glutathione S-transferase M1, glutathione S- | | | |
| M96233 s at | M96233 | 2433 | transferase M2 (muscle),glutathione S- | 4.23 | down | 0.04227 |
| M96843 at | M96843 | 2435 | EST | 8.42 | down | 0.02394 |
| M96843 at | M96843 | 2435 | EST | 4.08 | down | 0.02912 |
| M99439_at | M99439 | 2438 | transducin-like enhancer of split 4, homolog of | 5.14 | down | 0.00001 |
| rc N20113 s at | N20113 | 2439 | EST | 5.24 | down | 0.01346 |
| rc N21079 at | N21079 | 2441 | nucleolar cysteine-rich protein | 4.1 | down | 0.00028 |
| rc_N21550_at | N21550 | 2444 | EST | 3.08 | down | 0.00006 |
| rc N21646 at | N21646 | 2446 | EST | 3.79 | down | 0.00079 |
| rc_N22404_at | N22404 | 2450 | EST | 3.99 | down | 0.01152 |
| rc_N22434_at | N22434 | 2451 | EST | 4.37 | down | 0.01725 |
| rc_N22854_s_at | N22854 | 2452 | CASP2 and RIPK1 domain containing adaptor | 3.34 | down | 0.0084 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affry ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|----------|--------|--|-------------|-----------|---------|
| rc N22938 s at | · N22938 | 2453 | serum amyloid A4, constitutive | 35.39 | down | 0.00128 |
| rc N23665 s at | N23665 | 2455 | hydroxysteroid (17-beta) dehydrogenase 2 | 9.4 | down | 0.00055 |
| rc_N23730_s_at | N23730 | 2456 | v-fos FBJ murine osteosarcoma viral oncogene | 4.38 | down | 0.04395 |
| rc N23761 at | N23761 | 2457 | DKFZP586G011 protein | 5.41 | down | 0.00448 |
| N23817 at | N23817 | 2458 | EST | 3.76 | down | 0.00288 |
| rc_N24879_at | N24879 | 2460 | EST | 9.44 | down | 0.00008 |
| rc N25082 s at | N25082 | 2463 | amplified in osteosarcoma | 5.19 | down | 0.00895 |
| rc_N25193_at | N25193 | 2464 | EST | 3.13 | down | 0.01955 |
| l I | | | protein kinase, cAMP-dependent, regulatory, type | | | |
| rc N25969 s at | N25969 | 2466 | I, alpha (tissue specific extinguisher 1) | 4.32 | down | 0.00304 |
| rc N26184 at | N26184 | 2467 | MYLE protein | 4.82 | down | 0.00056 |
| rc_N27524_at | N27524 | 2472 | EST | 3.13 | down | 0.00216 |
| rc_N27563_at | N27563 | 2473 | EST | 9.3 8.3 | down | 0.00021 |
| N27670 at | N27670 | 2474 | progesterone membrane binding protein | 6.15 | down | 0.00321 |
| rc_N27834_at | N27834 | 2475 | alpha2,3-sialyltransferase | 5.31 | down | 0.00039 |
| rc_N29319_at | N29319 | 2476 | EST | 4.58 | down | 0.00011 |
| rc_N29353_at | N29353 | 2477 | kynurenine 3-monooxygenase (kynurenine 3- | 4.78 | down | 0.00019 |
| rc_N30856_at | N30856 | 2485 | solute carrier family 19 (thiamine transporter), | 3.71 | down | 0.00393 |
| rc_N31598_at | N31598 | 2488 | EST | 3.1 | down | 0.00203 |
| rc_N31741_at | N31741 | 2489 | serine hydroxymethyltransferase 1 (soluble) | 14.76 | down | 0.00001 |
| rc_N31952_at | N31952 | 2490 | EST | 3.13 | down | 0.01481 |
| rc_N32071_at | N32071 | 2491 | EST | 8.75 | down | 0.00006 |
| rc_N33009_s_at | N33009 | 2492 | apolipoprotein E | 60.54 | down | 0:0093 |
| rc N34441 at | N34441 | 2496 | EST | 3.07 | down | 0.00186 |
| rc_N34804_at | N34804 | 2497 | DKFZP434J214 protein | 8.08 | down | 0.00028 |
| rc_N36001_at | N36001 | 2504 | EST | 6.16 | down | 0.00222 |
| rc_N36250_at | N36250 | 2506 | cellular repressor of E1A-stimulated genes | 4.06 | down | 0.00776 |
| rc_N39163_at | N39163 | 2509 | metallothionein 1L | 4.3 | down | 0.03917 |
| rc_N39201_at | N39201 | 2510 | protease inhibitor 4 (kallistatin) | 24.91 | down | 0.00253 |
| rc_N40188_at | N40188 | 2513 | EST | 3.72 | down | 0.01771 |
| N40320_at | N40320 | 2514 | EST | 7.56 | down | 0.01584 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|-----------|---------|
| rc N45232 at | N45232 | 2517 | EST | 3.61 | down | 0.01308 |
| rc N45307 s at | N45307 | 2518 | EST | 4.55 | down | 0.00006 |
| rc N45998 at | N45998 | 2520 | EST | 3.14 | down | 0.00337 |
| rc_N47469_at | N47469 | 2522 | EST | 3.34 | down | 0.00329 |
| rc_N47942_at | N47942 | 2523 | progesterone membrane binding protein | 4.51 | down | 0.00168 |
| rc_N48180_at | N48180 | 2526 | EST | 3.36 | down | 0.00543 |
| rc_N48315_at | N48315 | 2527 | adaptor-related protein complex 2, mu 1 subunit | 5.3 | down | 0.0149 |
| rc_N48602_at | N48602 | 2529 | EST | 3.17 | down | 0.02913 |
| rc_N48674_at | N48674 | 2530 | EST | 4.06 | down | 0.00028 |
| rc_N48787_at | N48787 | 2531 | protease inhibitor 1 (anti-elastase), alpha-1- | 4.4 | down | 0.00292 |
| rc_N49090_at | N49090 | 2533 | EST | 18 | down | 0.00501 |
| rc N49104 s at | N49104 | 2534 | nuclear receptor interacting protein 1 | 3.83 | down | 0.00144 |
| rc N49113 at | N49113 | 2535 | EST | 3.4 | down | 0.00162 |
| rc_N49214_at | N49214 | 2536 | EST | 4.74 | down | 0.00064 |
| rc_N49595_at | N49595 | 2538 | EST | 10.39 | down | 0.00022 |
| rc_N49902_at | N49902 | 2540 | EST | 3.55 | down | 0.00455 |
| rc_N51117_at | N51117 | 2544 | EST | 89.6 | down | 0.00081 |
| rc_N51737_at | N51737 | 2547 | mitogen-activated protein kinase kinase | 3.32 | down | 0.00376 |
| rc_N51773_at | N51773 | 2549 | EST | 16.32 | down | 0.0007 |
| rc_N52271_at | N52271 | 2552 | LIM protein (similar to rat protein kinase C- | 8.06 | down | 0.00011 |
| rc_N52322_at | N52322 | 2553 | EST | 3.27 | down | 0.00933 |
| rc_N52845_at | N52845 | 2554 | EST | 5.53 | down | 0.00088 |
| rc_N52985_at | N52985 | 2555 | nidogen (enactin) | 4.21 | down | 0.01385 |
| rc N53031 s at | N53031 | 2556 | UDP glycosyltransferase 2 family, polypeptide B4 | 97.58 | down | 0.00022 |
| rc N53352 at | N53352 | 2558 | EST | 3.22 | down | 0.00416 |
| 1 | | | cytochrome P450, subfamily IIJ (arachidonic acid | | | |
| rc N53549 s at | N53549 | 2559 | epoxygenase) polypeptide 2 | 4.68 | down | 0.00818 |
| rc N53757 at | N53757 | 2560 | EST | 3.97 | down | 0.00255 |
| rc N54053 at | N54053 | 2561 | secreted phosphoprotein 2, 24kD | 60.39 | down | 0.00087 |
| rc_N54311_at | N54311 | 2564 | EST | 4.82 | down | 0.00183 |
| rc_N54399_at | N54399 | 2566 | EST | 3.34 | down | 0.00048 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affv ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|-----------|---------|
| rc N54417 s at | N54417 | 2567 | fibrinogen, A alpha polypeptide | 99.28 | down | 0.00001 |
| rc N54429 at | N54429 | 2568 | EST | 57.81 | down | 0.00724 |
| rc N54511 s at | N54511 | 2569 | KIAA0265 protein | 3.45 | down | 0.03362 |
| rc N54604 at | N54604 | 2570 | EST | 3.72 | down | 0.00741 |
| 1 | | | cytidine monophosphate-N-acetylneuraminic acid | | | |
| rc N54792 at | N54792 | 2571 | hydroxylase (CMP-N-acetylneuraminate | 3.23 | down | 0.00099 |
| rc N54950 s at | N54950 | 2573 | ketohexokinase (fructokinase) | 17.17 | down | 0.00078 |
| N57464 at | N57464 | 2576 | CCAAT/enhancer binding protein (C/EBP), delta | 14.69 | down | 0.00018 |
| rc N57934 s at | N57934 | 2577 | formiminotransferase cyclodeaminase | 13.81 | down | 0.00171 |
| rc N58326 at | N58326 | 2579 | EST | 14.97 | down | 0.00647 |
| rc_N59089_at | N59089 | 2581 | EST | 4.74 | down | 0.00055 |
| rc N59231 s at | N59231 | 2582 | pyruvate carboxylase | 3.45 | down | 0.02066 |
| rc_N59283_at | N59283 | 2583 | EST | 4.65 | down | 0.02343 |
| rc_N59474_at | N59474 | 2584 | EST | 6.93 | down | 0.00337 |
| rc N59532 s at | N59532 | 2585 | aminomethyltransferase (glycine cleavage | 6.73 | down | 0.00005 |
| rc_N59543_at | N59543 | 2587 | PDZ domain containing 1 | 96.6 | down | 0.00052 |
| rc_N59550_at | N59550 | 2588 | EST | 25.56 | down | 0.00024 |
| rc_N62443_at | N62443 | 2590 | EST | 3.72 | down | 0.01717 |
| rc_N62523_at | N62523 | 2592 | hepatic leukemia factor | 5.02 | down | 0.00087 |
| rc N62652 s at | N62652 | 2593 | EST | 5.23 | down | 0.03006 |
| rc N63391 at | N63391 | 2600 | EST | 7.39 | down | 0.00565 |
| rc N63688 at | N63688 | 2603 | EST | 3.13 | down | 0.002 |
| rc N63698 at | N63698 | 2604 | EST | 7.92 | down | 0.00001 |
| rc N63845 at | N63845 | 2605 | phytanoyi-CoA hydroxylase (Refsum disease) | 25.92 | down | 0.00084 |
| rc_N64017_at | N64017 | 2606 | EST | 5.27 | down | 0.00022 |
| | | | enoyl-Coenzyme A, hydratase/3-hydroxyacyl | | | • |
| rc N64036 s at | N64036 | 2607 | Coenzyme A dehydrogenase | 66.6 | down | 0.00125 |
| rc N64436 at | N64436 | 2609 | EST | 3.51 | down | 0.00441 |
| rc_N64535_at | N64535 | 2610 | EST | 6.19 | down | 0.00106 |
| rc_N65959_at | N65959 | 2612 | EST | 5.43 | down | 0.00044 |
| rc_N66066_at | 99099N | 2613 | EST | 28.35 | down | 0.00055 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| rc N66130 at | N66130 | 2614 | progesterone membrane binding protein | 3.98 | down | 0.0106 |
| rc_N66422_at | N66422 | 2616 | EST | 4.1 | down | 0.00237 |
| rc_N66763_at | N66763 | 2619 | EST | 3.79 | down | 0.03015 |
| rc_N66857_at | N66857 | 2620 | EST | 15.03 | down | 0.00005 |
| I | | | prion protein (p27-30) (Creutzfeld-Jakob disease, | | | |
| rc N67009 s at | 0029N | 2622 | Gerstmann-Strausler-Scheinker syndrome, fatal | 3.55 | down | 0.00956 |
| rc N67096 at | 96029N | 2623 | EST | 3.23 | down | 0.00446 |
| rc_N67105_at | N67105 | 2624 | EST | 5.01 | down | 0.00176 |
| rc_N67378_at | N67378 | 2626 | KIAA1053 protein | 7.14 | down | 0 |
| rc N67876 s at | N67876 | 2628 | insulin-like growth factor 1 (somatomedin C) | 8.89 | down | 0.00042 |
| rc N67893 at | N67893 | 2629 | EST | 10.48 | down | 0.00341 |
| rc N68596 s at | N68596 | 2636 | betaine-homocysteine methyltransferase | 40.99 | down | 0 |
| rc N68730 at | N68730 | 2637 | EST | 3.88 | down | 0.00091 |
| rc_N68974_at | N68974 | 2640 | EST | 16.44 | пмор | 0.00087 |
| rc_N68993_at | N68993 | 2641 | EST | 3.06 | down | 0.00867 |
| rc N69136 at | N69136 | 2645 | EST | 18.53 | . uwop | 0.00366 |
| rc_N69216_at | N69216 | 2646 | EST | 3.4 | down | 0.00497 |
| rc_N69299_at | N69299 | 2649 | EST | 3.63 | down | 0.03776 |
| rc_N70005_at | N70005 | 2653 | EST | 3.42 | down | 0.04229 |
| rc N70057 s at | N70057 | 2654 | DNA segment on chromosome 6 (unique) 49 | 5.24 | down | 0.01178 |
| rc_N70305_at | N70305 | 2655 | EST | 4.41 | down | 0.00078 |
| rc N70358 s at | N70358 | 2657 | growth hormone receptor | 34.35 | down | 0 |
| rc_N70861_at | N70861 | 2661 | EST | 6.55 | down | 0.00001 |
| l I | | | solute carrier family 10 (sodium/bile acid | | | |
| rc N70966 s at | 9960ZN | 2663 | cotransporter family), member 1 | 34.06 | down | 9000.0 |
| rc N71542 at | N71542 | 2665 | kidney- and liver-specific gene | 21.05 | down | 0 |
| rc n71935 s at | N71935 | 2667 | multiple PDZ domain protein | 16.28 | down | 0 |
| rc N72259 at | N72259 | 2670 | cornichon-like | 4.65 | down | 0.0054 |
| rc N72695 s at | N72695 | 2671 | EST | 9.18 | down | 0.00069 |
| , — | N73461 | 2673 | EST | 4.25 | down | 0.00162 |
| 1 ", | N73468 | 2674 | protein S (alpha) | 4.4 | down | 0.00853 |
| | | | | | | |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|-----------|---------|
| rc N73543 at | N73543 | 2675 | EST | 17 | down | 0.00003 |
| rc_N73561_at | N73561 | 2676 | EST | 12.96 | down | 0.00011 |
| rc_N73883_at | N73883 | 2682 | EST | 9.35 | down | 0 |
| rc_N73988_at | N73988 | 2683 | EST | 10.27 | down | 0.00083 |
| rc_N74025_at | N74025 | 2685 | deiodinase, iodothyronine, type I | 22.79 | down | 0 |
| rc_N74422_at | N74422 | 2686 | EST | 30.32 | down | 0 |
| rc_N74558_at | N74558 | 2687 | EST | 3.19 | down | 0.00247 |
| rc N75072 at | N75072 | 2689 | EST | 3.73 | down | 0.00379 |
| N75203 s at | N75203 | 2691 | EST | 6.15 | down | 0.00181 |
| N75870 s at | N75870 | 2693 | dual specificity phosphatase 1 | 13.41 | down | 0.00251 |
| rc N76012 r at | N76012 | 2694 | EST | 36.71 | down | 0.00598 |
| N77326 at | N77326 | 2696 | EST | 7.51 | down | 0.00542 |
| N77606_at | N77606 | 2697 | EST | 4.22 | down | 0.00119 |
| ļ | | | macrophage stimulating 1 (hepatocyte growth | | | |
| rc N78850 s at | N78850 | 2699 | factor-like), macrophage stimulating, pseudogene | 13.69 | down | 0.00421 |
| rc N78902 at | N78902 | 2700 | leptin receptor | 6.79 | down | 0.0041 |
| rc_N79435_at | N79435 | 2701 | chromosome 15 open reading frame 3 | 3.12 | down | 0.00861 |
| rc_N79778_at | N79778 | 2702 | extracellular matrix protein 2, female organ and | 5.15 | down | 0.00286 |
| rc_N80129_i_at | N80129 | 2703 | metallothionein 1L | 66.48 | down | 0.00415 |
| rc_N80129_f_at | N80129 | 2703 | metallothionein 1L | 13.6 | down | 0.00196 |
| rc N81025 at | N81025 | 2705 | EST | 8.61 | down | 0.00015 |
| rc_N81036_at | N81036 | 2706 | EST | 68.9 | down | 0.00276 |
| N89302 s at | N89302 | 2708 | HLA-B associated transcript-3 | 23.44 | down | 0.00192 |
| rc N89738 at | N89738 | 2710 | EST | 10.06 | down | 0.00052 |
| rc_N90584_at | N90584 | 2714 | EST | 3.08 | down | 0.02602 |
| N90820 at | N90820 | 2715 | DKFZP566B1346 protein | 4.86 | down | 0.03008 |
| N91087_at | N91087 | 2717 | EST | 6.91 | down . | 0.00109 |
| rc N91273 r at | N91273 | 2718 | EST | 4.07 | down | 0.02965 |
| rc N91882 at | N91882 | 2720 | alpha2,3-sialyltransferase | 14.52 | down | 0.00024 |
| rc_N93155 f_at | N93155 | 2729 | calmodulin 1 (phosphorylase kinase, delta) | 4.39 | down | 0.0002 |
| rc_N93155_i_at | N93155 | 2729 | calmodulin 1 (phosphorylase kinase, delta) | 3.46 | down | 0.00031 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affv ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------------|---------|--------|--|-------------|-----------|---------|
| rc N93191 at | N93191 | 2730 | EST | 3.25 | down | 0.00232 |
| rc N93246 f at | N93246 | 2731 | EST | 3.03 | down | 0.00049 |
| rc N93764 at | N93764 | 2737 | EST | . 5.92 | down | 0.0109 |
| N94146 at | N94146 | 2739 | EST | 8.05 | down | 0 |
| rc N94367 at | N94367 | 2740 | EST | 8.79 | down | 0.01003 |
| rc_N94930_at | N94930 | 2741 | multiple PDZ domain protein | 10.06 | down | 0.00226 |
| rc N95495 at | N95495 | 2742 | EST | 5.61 | down | 0.00308 |
| rc_N95585_at | N95585 | 2743 | EST | 3.1 | down | 0.0083 |
| N99542 at | N99542 | 2747 | orosomucoid 1 | 8.41 | down | 0.00001 |
| rc N99866 at | 99866N | 2748 | EST | 3.13 | down | 0.04148 |
| rc_R00296_at | R00296 | 2750 | EST | 3.98 | down | 0.04632 |
| rc R00843 s at | R00843 | 2751 | fragile histidine triad gene | 3.68 | down | 0.03578 |
| rc_R01023_s_at | R01023 | 2752 | glucokinase (hexokinase 4) regulatory protein | 50.71 | down | 0.00321 |
| rc R01081 at | R01081 | 2753 | EST | 3.3 | down | 0.00839 |
| ! ! | | | glucose-6-phosphatase, catalytic (glycogen | | | |
| rc R02365 f at | R02365 | 2755 | storage disease type I, von Gierke disease) | 17.17 | down | 0.00124 |
| rc R02371 at | R02371 | 2756 | EST | 3.75 | down | 60000.0 |
| rc_R02572_at | R02572 | 2757 | fibronectin 1 | 7.1 | down | 0.00059 |
| rc_R02752_at | R02752 | 2758 | EST | 3.05 | down | 0.00362 |
| rc <u>R05309 f</u> at | R05309 | 2759 | EST | 3.72 | down | 0.0008 |
| rc R05490 at | R05490 | 2761 | SEC24 (S. cerevisiae) related gene family, | 3.03 | down | 0.00317 |
| rc_R05518_at | R05518 | 2762 | EST | 4.15 | down | 0.0125 |
| rc <u>R</u> 06002 s at | R06002 | 2763 | EST | 10.15 | down | 0.00003 |
| rc_R06271_s_at | R06271 | 2766 | EST | 6.14 | down | 0.00063 |
| rc_R06726_s_at | R06726 | 2770 | protease inhibitor 1 (anti-elastase), alpha-1- | 12.26 | down | 0.03905 |
| rc R06746 at | R06746 | 2771 | EST | 15.77 | down | 60000.0 |
| rc R06764 s at | R06764 | 2772 | apolipoprotein B (including Ag(x) antigen) | 4.91 | down | 0.00167 |
| rc R06977 f at | R06977 | 2775 | glucokinase (hexokinase 4) regulatory protein | 6.1 | down | 0.00049 |
| rc R07637 at | R07637 | 2778 | EST | 3.04 | down | 0.00118 |
| | R08548 | 2779 | | 9.94 | down | 0.00326 |
| rc R08564 at | R08564 | 2780 | plasminogen-like | 60.18 | down | 0.00091 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|-----------|---------|
| rc R08615 s at | R08615 | 2781 | homogentisate 1,2-dioxygenase (homogentisate | 21.85 | down | 0.00026 |
| rc R08850 at | R08850 | 2782 | EST | 12.55 | down | 0.0000 |
| rc_R09053_at | R09053 | 2783 | EST | 5.9 | down | 0.0002 |
| rc_R10138_at | R10138 | 2784 | EST | 4 | down | 0.00032 |
| rc_R10287_at | R10287 | 2785 | EST | 11.2 | down | 0.0003 |
| rc R10378 s at | R10378 | 2786 | fibrinogen-like 1 | 8.13 | down | 0.00189 |
| rc_R10662_f at | R10662 | 2787 | mutL (E. coli) homolog 1 (colon cancer, | 3.2 | down | 0.0005 |
| rc R10684 at | R10684 | 2788 | EST | 5.51 | down | 0.00741 |
| rc_R12472_at | R12472 | 2789 | EST | 55.18 | down | 0.00011 |
| rc_R12579_at | R12579 | 2790 | EST | 3.5 | down | 0.00137 |
| rc R15825 r at | R15825 | 2792 | KIAA0946 protein; Huntingtin interacting protein | 5.33 | down | 0.00391 |
| rc_R16098_at | R16098 | 2793 | EST | 63.41 | down | 0.00038 |
| rc_R17762_at | R17762 | 2795 | EST | 3.37 | down | 0.01822 |
| rc_R21232_at | R21232 | 2798 | EST | 3.56 | down | 0.00246 |
| rc_R22196_at | R22196 | 2799 | EST · | 3.75 | down | 0.02867 |
| rc_R22905_at | R22905 | 2801 | EST | 4.64 | down | 0.0043 |
| rc_R26904 f_at | R26904 | 2805 | EST | 3.51 | down | 0.00058 |
| R31641_at | R31641 | 2814 | EST | 96.6 | down | 0.00011 |
| rc_R31917_s_at | R31917 | 2815 | EST | 3.91 | down | 0.00071 |
| R3203 | R32036 | 2816 | interleukin 1 receptor-like 1 | 4.02 | down | 0.00051 |
| R3244 | R32440 | 2817 | EST · | 7.41 | down | 0.00159 |
| rc R32490 s at | R32490 | 2818 | EST | 8.95 | down | 0.00215 |
| rc_R33146_at | R33146 | 2819 | EST | 7 | down | 0.00043 |
| rc R34133 at | R34133 | 2821 | EST | 4.13 | down | 0.00008 |
| rc_R34362_at | R34362 | 2822 | KIAA0327 gene product | 5.4 | down | 0.04615 |
| rc_R36228_at | R36228 | 2824 | EST | 4.39 | down | 0.00033 |
| rc R36989 s at | R36989 | 2826 | hypothetical protein, estradiol-induced | 13.7 | down | 0.00056 |
| rc R37128 s at | R37128 | 2827 | complement component 4A | 39.51 | down | 0.00364 |
| rc R38185 at | R38185 | 2829 | EST | 6.32 | down | 0.00063 |
| rc_R38709_s_at | R38709 | 2833 | superoxide dismutase 2, mitochondrial | 8.23 | down | 0.01578 |
| rc_R39234_r_at | R39234 | 2835 | EST | 3.24 | down | 0.0412 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change Direction | Direction | Pvalue |
|------------------------|---------|--------|--|-----------------------|-----------|---------|
| rc R40395 s at | R40395 | 2841 | lecithin-cholesterol acyltransferase | 37.33 | down | 0.00032 |
| rc R40492 at | R40492 | 2842 | EST | 8.89 | down | 0.00229 |
| rc <u>F</u> 40556 s at | R40556 | 2843 | EST | 3.57 | down | 0.00184 |
| rc R40899 f at | R40899 | 2844 | glycine receptor, beta | 9.11 | down | 6000.0 |
| rc_R40946 f at | R40946 | 2845 | crystallin, zeta (quinone reductase) | 6.14 | down | 0.00156 |
| rc R42241 at | R42241 | 2846 | EST | 3.97 | down | 0.00129 |
| rc R43166 i at | R43166 | 2847 | EST | 3.37 | down | 0.00919 |
| rc_R43174 s at | R43174 | 2848 | paraoxonase 1 | 74.04 | down | 0.00038 |
| rc R43365 at | R43365 | 2850 | EST | 5.37 | down | 0.00103 |
| rc_R43799_at | R43799 | 2851 | EST | 16.35 | down | 0.00208 |
| rc R43910 at | R43910 | 2852 | EST | 5.71 | down | 0 |
| rc_R44025_at | R44025 | 2854 | EST | 4.4 | down | 0.01325 |
| rc_R44761_at | R44761 | 2858 | aryl hydrocarbon receptor nuclear translocator | 3.12 | down | 0.02663 |
| rc <u>R</u> 45656 i at | R45656 | 2865 | EST | 10.21 | down | 0.00179 |
| rc R48307 at | R48307 | 2870 | EST | 7.18 | down | 0.00007 |
| rc R48540 s at | R48540 | 2873 | EST | 6.05 | down | 0.00086 |
| rc R48732 s at | R48732 | 2876 | EST | 5.12 | down | 0.00837 |
| rc R49035 at | R49035 | 2877 | EST | 8.02 | down | 0.00991 |
| rc <u>R49459 s</u> at | R49459 | 2882 | transferrin receptor 2 | 85.61 | down | 0.00048 |
| rc R49602 at | R49602 | 2885 | EST | 36.5 | down | 0.00001 |
| R50008 s at | R50008 | 2887 | 7-dehydrocholesterol reductase | 29.9 | down | 0.00409 |
| rc R51256 at | R51256 | 2889 | EST | 3.04 | down | 0.00286 |
| rc_R51309_at | R51309 | 2890 | KIAA1077 protein | 3.54 | down | 0.04112 |
| rc_R51831_at | R51831 | 2891 | EST | 5.03 | down | 0.00761 |
| rc_R52800_at | R52800 | 2895 | EST | 3.96 | down | 0.00749 |
| rc R52822 i at | R52822 | 2896 | EST | 12.79 | down | 0.00001 |
| rc R52949 at | R52949 | 2897 | EST | 3.46 | down | 0.00395 |
| rc R54416 at | R54416 | . 2901 | EST | 3.14 | down | 0.00236 |
| rc_R56094_at | R56094 | 2905 | EST | 4.87 | down | 0.00214 |
| rc_R56602_at | R56602 | 2907 | lg superfamily protein | 3.06 | down | 0.02464 |
| rc_R59221_at | R59221 | 2912 | progesterone binding protein | 7.54 | down | 0.00159 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|---------|--------|---|-------------|-----------|---------|
| rc R59325 at | R59325 | 2914 | EST | 4.48 | down | 0.00117 |
| rc R59722 at | R59722 | 2916 | EST | 19.74 | down | 0.00016 |
| rc R61740 f at | R61740 | 2923 | EST | 4.02 | down | 0.00321 |
| rc R62173 f at | R62173 | 2924 | UDP-glucose dehydrogenase | 5.76 | down | 9000'0 |
| rc R62519 f at | R62519 | 2926 | EST | 10.83 | down | 0.00243 |
| R63545 at | R63545 | 2927 | EST | 5.4 | down | 0.00202 |
| rc R64131 at | R64131 | 2930 | EST | 20.58 | down | 0.01028 |
| rc R64144 s at | R64144 | 2932 | cAMP responsive element binding protein-like 2 | 4.58 | down | 0.00495 |
| | R64199 | 2933 | SEC22, vesicle trafficking protein (S. cerevisiae)- | 9.19 | down | 0.00387 |
| rc R65593 s at | R65593 | 2935 | kynurenine 3-monooxygenase (kynurenine 3- | 21.27 | down | 0.00007 |
| R67751 at | R67751 | 2939 | EST | 3.12 | down | 0.00319 |
| rc R67970 s at | R67970 | 2940 | gamma-glutamyl carboxylase | 9.45 | down | 0.00212 |
| rc R69031 at | R69031 | 2941 | EST | 3.19 | down | 0.00798 |
| R69417 at | R69417 | 2942 | EST | 7.79 | down | 0.03795 |
| rc_R70790 at | R70790 | 2948 | EST | 4.07 | down | 0.02354 |
| rc R71491 at | R71491 | 2954 | EST | 3.55 | down | 0.01903 |
| rc_R73485_at | R73485 | 2958 | EST | 7.77 | down | 6000.0 |
| rc_R73816_at | R73816 | 2961 | EST . | 11.53 | down | 0.00259 |
| R77628 at | R77628 | 2966 | insulin induced gene 1 | 29.23 | down | 0.00122 |
| R80048 at | R80048 | 2972 | EST · | 12.28 | down | 0.00128 |
| R80573 at | R80573 | 2973 | EST | 3.08 | down | 0.02126 |
| rc_R82074 f at | R82074 | 2974 | syndecan 1 | 99.9 | down | 0.01336 |
| R82229 at | R82229 | 2975 | phosphatidylserine decarboxylase | 3.08 | down | 0.03455 |
| rc R82837 at | R82837 | 2976 | KIAA0970 protein | 5 | down | 0.00181 |
| R87373 s at | R87373 | 2978 | EST | 4.09 | down | 0.02253 |
| rc R89811 s at | R89811 | 2980 | HGF activator | 62.51 | down | 0.00148 |
| rc_R91503_s_at | R91503 | 2982 | ATP-binding cassette, sub-family C (CFTR/MRP), | 9.64 | down | 0.00584 |
| rc_R92458 f_at | R92458 | 2986 | hemoglobin, gamma G | 3.31 | down | 0.00763 |
| rc_R92475_s_at | R92475 | 2987 | flavin containing monooxygenase 3 | 26.13 | down | 0 |
| rc_R92737_at | R92737 | 2988 | EST | 4.53 | down | 0.04447 |
| rc_R92768_at | R92768 | 2989 | EST | 13.2 | down | 0.00001 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affv ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------------|---------|--------|--|-------------|-----------|---------|
| rc R93714 at | R93714 | 2992 | fetuin B | 10.42 | down | 0.00043 |
| R93776 s at | R93776 | 2993 | EST | 39.32 | down | 0.00176 |
| rc R93908 at | R93908 | 2994 | EST | 5.8 | down | 0.02699 |
| F94662 at | R94662 | 2995 | heme-binding protein | 4.41 | down | 0.02301 |
| rc R94674 s at | R94674 | 2996 | EST | 10.76 | down | 0.00008 |
| R96417 at | R96417 | 2998 | EST | 3.9 | down | 0.00019 |
| rc R9682 <u>2</u> at | R96822 | 3000 | EST | 8.25 | down | 0.00008 |
| rc_R97302_at | R97302 | 3003 | EST | 3.73 | down | 0.01887 |
| 1 | | | cytochrome P450, subfamily VIIIB (sterol 12- | | | |
| rc R97419 at | R97419 | 3004 | alpha-hydroxylase), polypeptide 1 | 65.07 | down | 0.0039 |
| rc_R97711_at | R97711 | 3005 | EST | 3.37 | down | 0.00838 |
| rc_R97798_at | R97798 | 3007 | EST | 3.79 | down | 0.00015 |
| E98073 at | R98073 | 3009 | EST | 46.87 | down | O |
| rc R98074 at | R98074 | 3010 | EST | 9.21 | down | 0.00048 |
| rc_R98413_at | R98413 | 3012 | EST | 9.71 | down | 0.00007 |
| rc_R98624_at | R98624 | 3013 | EST | 21.32 | down | 0 |
| rc_R98774_at | R98774 | 3014 | EST | 3.93 | down | 0.00001 |
| rc_R99591_at | R99591 | 3016 | CD5 antigen-like (scavenger receptor cysteine | 9.52 | down | 0.00006 |
| rc_R99909_at | R99909 | 3017 | EST | 4.27 | down | 0.00045 |
| S48983 at | S48983 | 3018 | serum amyloid A4, constitutive | 23.04 | down | 0.00022 |
| S52028 s at | S52028 | 3019 | cystathionase (cystathionine gamma-lyase) | 3.21 | down | 0.00476 |
| S62539 s at | S62539 | 3022 | insulin receptor substrate 1 | 3.7 | down | 0.01307 |
| S67325 at | S67325 | 3024 | propionyl Coenzyme A carboxylase, beta | 4.19 | down | 0.00151 |
| I | | | aldo-keto reductase family 1, member C4 | | | |
| | | | (chlordecone reductase; 3-alpha hydroxysteroid | | | |
| S68287 at | S68287 | 3025 | dehydrogenase, type I; dihydrodiol | 24.11 | down | 0 |
| S68805 at | S68805 | 3026 | glycine amidinotransferase (L-arginine:glycine | 18.19 | down | 0.00083 |
| S69232 at | S69232 | 3027 | electron-transferring-flavoprotein dehydrogenase | 4.9 | down | 0.00017 |
| S70004 at | S70004 | 3029 | glycogen synthase 2 (liver) | 96.6 | down | 0.00001 |
| S72370_at | S72370 | 3030 | pyruvate carboxylase | 5.31 | down | 0.00075 |
| S72370_at | S72370 | 3030 | pyruvate carboxylase | 4.54 | down | 0.0001 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| S74728 at | S74728 | 3033 | antiquitin 1 | 5.01 | down | 0.00008 |
| S77356_at | S77356 | 3034 | EST | 5.83 | down | 0.00122 |
| S77410 at | S77410 | 3035 | angiotensin receptor 1 | 10.4 | down | 0 |
| S90469 at | S90469 | 3040 | P450 (cyfochrome) oxidoreductase | 4.32 | down | 0.0182 |
| S95936 at | S95936 | 3041 | transferrin | 19.76 | down | 0.02009 |
| rc T03441 f at | T03441 | 3044 | cytochrome b-561 | 5.74 | down | 0.02781 |
| rc_T03651_s_at | T03651 | 3047 | tubulin, beta polypeptide | 6.47 | down | 0.00843 |
| T08879 at | T08879 | 3049 | cathepsin F | 5.29 | down | 0.0008 |
| rc T10108 s at | T10108 | 3050 | EST | 3.85 | down | 0.01155 |
| rc 110264 s at | T10264 | 3051 | EST | 3.84 | down | 0.00874 |
| rc T10322 at | T10322 | 3053 | dihydropyrimidinase-like 2 | 4.23 | down | 0.01527 |
| rc_T10822_at | T10822 | 3055 | EST | 3.16 | down | 0.00635 |
| rc T15482 f at | T15482 | 3060 | EST | 5.04 | down | 0.00233 |
| rc_T15674 f at | T15674 | 3061 | EST | 4.12 | down | 0.02111 |
| rc T16175 s at | T16175 | 3064 | protein tyrosine phosphatase, non-receptor type | 3.07 | down | 0.01007 |
| rc T16269 at | T16269 | 3067 | EST | 3.36 | down | 0.00282 |
| rc_T16478_at | T16478 | 3070 | EST | 4 | down | 0.01041 |
| rc T16484 s at | T16484 | 3071 | EST | 69.7 | down | 0.00805 |
| rc T17411 s at | T17411 | 3078 | transthyretin (prealbumin, amyloidosis type I) | 13.49 | down | 0.00203 |
| rc T23430 at | T23430 | 3080 | EST | 3.12 | down | 0.00949 |
| rc_T23680_at | T23680 | 3084 | calcium channel, voltage-dependent, gamma | 3.29 | down | 0.0003 |
| rc T23882 s at | T23882 | 3085 | kininogen | 42.85 | down | 0.00641 |
| rc_T23986_s at | T23986 | 3086 | hydroxyacyl glutathione hydrolase; glyoxalase 2 | 7.26 | down | 0.0062 |
| rc T24055 s at | T24055 | 3087 | ribosomal protein L26 | 3.9 | down | 0.00046 |
| rc T24106 at | T24106 | 3089 | EST | 12.13 | down | 0.01687 |
| rc_T25506_at | T25506 | 3090 | EST | 4.51 | down | 0.00529 |
| T30341 s at | T30341 | 3101 | EST | 3.04 | down | 0.04567 |
| rc T33011 at | T33011 | 3104 | EST | 3.57 | down | 0.02486 |
| T39897 s at | T39897 | 3114 | androgen induced protein | 10 | down | 0.00466 |
| rc T40936 at | T40936 | 3118 | EST | 46.92 | down | 0.00056 |
| rc_T40995_f_at | T40995 | 3119 | alcohol dehydrogenase 3 (class I), gamma | 13.25 | down | 0.01531 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|-----------|---------|
| rc T41047 s at | T41047 | 3120 | EST | 4.74 | down | 0.00015 |
| rc T41232 at | T41232 | 3122 | EST | 3.57 | down | 0.00846 |
| rc_T46901_at | T46901 | 3123 | EST | 77.28 | down | 9000.0 |
| rc T47778 s at | T47778 | 3127 | fibrinogen, A alpha polypeptide | 30 | down | 0.01401 |
| rc T48039 s at | T48039 | 3129 | protein C (inactivator of coagulation factors Va | 84.39 | down | 0.00112 |
| rc_T48075_f_at | T48075 | 3130 | hemoglobin, alpha 1 | 32.56 | down | 0.00172 |
| rc T48278 at | T48278 | 3132 | EST | 15.04 | down | 0.04751 |
| rc T48980 s at | T48980 | 3134 | calmodulin 1 (phosphorylase kinase, delta) | 4.2 | down | 0.0046 |
| T49061 | T49061 | 3135 | EST | 3.06 | down | 0.00768 |
| rc_T51150_at | T51150 | 3137 | EST | 10.52 | down | 0.00377 |
| 1 | | | solute carrier family 22 (extraneuronal | | | |
| rc T51617 at | T51617 | 3138 | monoamine transporter), member 3 | 17.67 | down | 0.00018 |
| rc_T51930_at | T51930 | 3139 | EST | 66.6 | down | 0.00066 |
| rc_T52564_at | T52564 | 3141 | EST | 3.28 | down | 0.01506 |
| rc_T52813 s at | T52813 | 3142 | putative lymphocyte G0/G1 switch gene | 16.71 | down | 0.03897 |
| rc_T54160_s_at | T54160 | 3145 | EST | 3.24 | down | 0.00168 |
| rc_T55547_at | T55547 | 3148 | EST | 3.28 | down | 0.00047 |
| rc T56264 s at | T56264 | 3149 | apolipoprotein C-II | 53.04 | down | 0.00938 |
| rc_T56279_s_at | T56279 | 3150 | H factor (complement)-like 3 | 28.39 | down | 0.00016 |
| rc_T56281_f_at | T56281 | 3151 | RNA helicase-related protein | 32.34 | down | 0.00002 |
| T57140 s at | T57140 | 3152 | paraoxonase 3 | 28.8 | down | 0 |
| rc T58032 s at | T58032 | 3153 | 3-hydroxyanthranilate 3,4-dioxygenase | 8.86 | down | 0.00023 |
| rc_T58756_at | T58756 | 3156 | EST | 13.39 | down | 0.00013 |
| rc_T58775_at | T58775 | 3157 | small inducible cytokine subfamily A (Cys-Cys), | 21.18 | down | 900000 |
| rc T59148 s at | T59148 | 3158 | carbamoyf-phosphate synthetase 1, mitochondrial | 88.89 | down | 0 |
| rc T60407 at | T60407 | 3161 | EST | 9.9 | down | 0.00167 |
| rc_T61256_s_at | T61256 | 3162 | ketohexokinase (fructokinase) | 13.59 | down | 0.00425 |
|]] | | | vitronectin (serum spreading factor, somatomedin | | | • |
| rc_T61373_s_at | T61373 | 3163 | B, complement S-protein) | 44.9 | down | 0.03172 |
| rc_T61389 f_at | T61389 | 3164 | haptoglobin | 18.95 | down | 0.04873 |
| rc_T61649_f_at | T61649 | 3165 | superoxide dismutase 2, mitochondrial | 26.09 | down | 0.00734 |
| | | | | | | |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Genbank | Sed ID | Known Gene Name | Fold Change | | Pvalue |
|---------|--------|--|-------------|------|---------|
| 61801 | 3167 | hemopexin | 25.7 | down | 0.01362 |
| 63364 | 3170 | ficolin (collagen/fibrinogen domain-containing) 3 | 6.72 | down | 0.00383 |
| | 3171 | EST | 16.54 | down | 0 |
| | 3172 | EST | 10.46 | down | 0.00014 |
| | 3173 | protein phosphatase 5, catalytic subunit | 2.66 | down | 0.00349 |
| | 3177 | EST | 3.21 | down | 0.04847 |
| | 3178 | glutaryl-Coenzyme A dehydrogenase | 8.61 | down | 0.00003 |
| | | succinate dehydrogenase complex, subunit D, | | | |
| | 3181 | integral membrane protein | 7.04 | down | 0.03554 |
| | 3183 | asialoglycoprotein receptor 2 | 31.6 | down | 0.00705 |
| | 3184 | fibrinogen, B beta polypeptide | 49.55 | down | 0 |
| | 3185 | short-chain dehydrogenase/reductase 1 | 4.03 | down | 0.01593 |
| | 3186 | CD81 antigen (target of antiproliferative antibody | 3.11 | down | 0.01634 |
| | 3187 | EST | 3.87 | down | 0.00617 |
| | 3188 | EST | 37.65 | down | 0.00036 |
| | 3189 | EST | 34.04 | down | 0 |
| | 3190 | metallothionein 1L | 28.72 | down | 0.02953 |
| | 3191 | carboxylesterase 1 (monocyte/macrophage | 60.35 | down | 0.00409 |
| | 3192 | quinoid dihydropteridine reductase | 6.54 | down | 0.00145 |
| | 3193 | EST | 5.39 | down | 0.00383 |
| | 3194 | haptoglobin | 33.18 | down | 0.02825 |
| | 3195 | EST | 4.38 | down | 0.00548 |
| | 3196 | mannose-binding lectin (protein C) 2, soluble | 36.53 | down | 0 |
| | 3198 | period (Drosophila) homolog 1 | 5.88 | down | 0.01219 |
| | 3199 | heat shock 90kD protein 1, beta | 3.98 | down | 0.00786 |
| | 3200 | EST | 4.7 | down | 0.00173 |
| | 3201 | fibrinogen, B beta polypeptide | 43.61 | down | 0.00743 |
| | 3202 | EST | 6.74 | down | 0.0017 |
| | 3203 | EST | 83.08 | down | 0.00069 |
| | 3204 | EST | 10.3 | down | 0.00897 |
| | 3205 | EST | 4.39 | down | 0.0017 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|-----------|---------|
| rc T72171 s at | T72171 | 3206 | thyroxin-binding globulin | 10.41 | down | 0.00163 |
| rc_T72268_s_at | T72268 | 3207 | B-factor, properdin | 7.43 | down | 0.01197 |
| rc T72502 at | T72502 | 3208 | EST | 8.87 | down | 0.0000 |
| rc T72629 i at | T72629 | 3209 | EST | 3.24 | down | 0.00556 |
| rc T72906 at | T72906 | 3210 | EST | 7.56 | down | 0.00062 |
| rc_T73420_at | T73420 | 3211 | short-chain dehydrogenase/reductase 1 | 4.58 | down | 0.00656 |
| rc T73433 s at | T73433 | 3212 | angiotensinogen | 48.3 | down | 0.00049 |
| rc T73442 at | T73442 | 3213 | EST | 94.41 | down | 0 |
| l [*] | | | alanine-glyoxylate aminotransferase (oxalosis I; | | | |
| rc T73739 s at | T73739 | 3214 | hyperoxaluria I; glycolicaciduria; serine-pyruvate | 16.18 | down | 0.00277 |
| rc T74542 s at | T74542 | 3215 | UDP glycosyltransferase 2 family, polypeptide | 32.36 | down | 0.00004 |
| rc T74608 at | T74608 | 3216 | hydroxyacid oxidase (glycolate oxidase) 1 | 14.03 | down | 0 |
| rc T74884 r at | T74884 | 3217 | EST | 3.76 | down | 0.02547 |
| rc_T77729 f at | T77729 | 3218 | pyruvate carboxylase | 7.29 | down | 0.00022 |
| יט | T78433 | 3220 | glycogen synthase 2 (liver) | 8.62 | down | 0.00072 |
| ွလ | T78889 | 3221 | fibronectin 1 | 19.87 | down | 0.00912 |
| rc_T79758_at | T79758 | 3224 | EST | 3.94 | down | 0.00581 |
| rc T79842 at | T79842 | 3225 | EST | 3.16 | down | 0.03159 |
| rc_T79863_at | T79863 | 3226 | EST | 5.62 | down | 0.00074 |
| rc_T81315_at | T81315 | 3227 | EST | 4.28 | down | 0.00187 |
| rc_T82254_at | T82254 | 3229 | EST | 3.82 | down | 0.00091 |
| rc_T82259_at | T82259 | 3230 | EST | 3.12 | down | 0.01122 |
| rc T82323 at | T82323 | 3231 | immunoglobulin superfamily, member 4 | 10.14 | down | 0 |
| rc T83356 s at | T83356 | 3232 | apolipoprotein H (beta-2-glycoprotein I) | 64.34 | down | 0.00802 |
| T83397 at | T83397 | 3233 | phytanoyl-CoA hydroxylase (Refsum disease) | 63.6 | down | 0 |
| T85532 f at | T85532 | 3236 | EST | 4.33 | down | 0.00591 |
| rc_T86464_at | T86464 | 3237 | EST | 3.17 | down | 0.00498 |
| rc_T86482_at | T86482 | 3238 | transferrin | 11.15 | down | 0.00006 |
| rc T86978 s at | T86978 | 3239 | glutamate dehydrogenase 1 | 9.44 | down | 0.00006 |
| rc_T87174_at | T87174 | 3240 | EST | 21.71 | down | 0.00681 |
| rc_T90037_at | T90037 | 3246 | EST | 3.99 | down | 0.0016 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change Direction | Direction | Pvalue |
|----------------|---------|--------|--|-----------------------|-----------|---------|
| rc T90492 at | T90492 | 3248 | EST | 3.88 | down | 0.00454 |
| rc_T90520_at | T90520 | 3249 | EST | 8.67 | down | 0.00072 |
| rc_T90531_at | T90531 | 3250 | EST | 3.6 | down | 0.00301 |
| rc_T90841_f at | T90841 | 3251 | EST | 5.74 | down | 0.01876 |
| rc_T91161_at | T91161 | 3253 | EST | 3.48 | down | 0.00002 |
| rc_T91348_at | T91348 | 3254 | EST | 3.57 | down | 0.00011 |
| rc_T92950_at | T92950 | 3256 | EST | 3.01 | down | 0.00497 |
| rc_T94862_at | T94862 | 3258 | EST | 3.07 | down | 0.01417 |
| rc_T95064_at | T95064 | 3260 | EST | 5.75 | down | 0.00604 |
| rc_T95515_s at | T95515 | 3261 | KIAA0249 gene product | 6.77 | down | 0.0001 |
| T95813 f at | T95813 | 3262 | KIAA1051 protein | 38.38 | down | 0.00008 |
| rc_T96969_at | T96969 | 3264 | EST | 3.42 | down | 0.00053 |
| rc_T97234_at | T97234 | 3265 | EST | 4.34 | down | 0.00263 |
| rc_T98199 i at | T98199 | 3267 | EST | 4.86 | down | 0.00431 |
| rc_T98676_at | T98676 | 3269 | EST | 46.94 | down | 0.0001 |
| rc T99636 s at | T99636 | 3271 | complement component 3 | 29.6 | down | 0.00051 |
| U00115_at | U00115 | 3272 | B-cell CLL/lymphoma 6 (zinc finger protein 51) | 5.18 | down | 0.00045 |
| | | | glucose-6-phosphatase, catalytic (glycogen | - | | |
| U01120_at | U01120 | 3274 | storage disease type I, von Gierke disease) | 13.41 | down | 0.00147 |
| • | | | solute carrier family 1 (glial high affinity glutamate | | | |
| U01824_at | U01824 | 3276 | transporter), member 2 | 3.2 | down | 0.00021 |
| U02020_at | U02020 | 3277 | pre-B-cell colony-enhancing factor | 3.37 | down | 0.01112 |
| U02388_at | U02388 | 3278 | cytochrome P450, subfamily IVF, polypeptide 2 | 9.38 | down | 0.00001 |
| U03056_at | 03056 | 3280 | hyaluronoglucosaminidase 1 | 6.64 | down | 0 |
| U03105_at | U03105 | 3281 | proline-rich protein with nuclear targeting signal | 4.69 | down | 0.00017 |
| | | | aldo-keto reductase family 1, member C1 | | | |
| | | | (dihydrodiol dehydrogenase 1; 20-alpha (3-alpha)- | | | |
| U05861_at | U05861 | 3285 | hydroxysteroid dehydrogenase) | 23.22 | down | 0.00002 |
| U06641_s_at | U06641 | 3287 | UDP glycosyltransferase 2 family, polypeptide | 24.32 | down | 0.00001 |
| U06863_at | U06863 | 3288 | follistatin-like 1 | 3.09 | down | 0.00091 |
| U08006_s_at | 00800 | 3290 | complement component 8, alpha polypeptide | 16 | down | 0.00111 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|-----------|---------|
| U08198_rna1_at | U08198 | 3292 | complement component 8, gamma polypeptide | 19.71 | down | 0 |
| U08854 s at | U08854 | 3293 | UDP glycosyltransferase 2 family, polypeptide | 7.38 | down | 0.00005 |
| U11313_at | U11313 | 3297 | sterol carrier protein 2 | 5.44 | down | 0.00187 |
| U12778_at | U12778 | 3301 | acyl-Coenzyme A dehydrogenase, | 5.68 | down | 0.00116 |
| | | | sulfotransferase family 2A, | | | |
| U13061_rna1_at | U13061 | 3302 | dehydroepiandrosterone (DHEA) -preferring, | 11.32 | down | 0.00048 |
| U16660 at | U16660 | 3313 | enoyl Coenzyme A hydratase 1, peroxisomal | 3.96 | down | 0.00002 |
| U19495 s at | U19495 | 3319 | stromal cell-derived factor 1 | 4.59 | down | 0.00011 |
| U19523 at | U19523 | 3320 | GTP cyclohydrolase 1 (dopa-responsive | 5.25 | down | 0.00029 |
| U20530_at | U20530 | 3322 | secreted phosphoprotein 2, 24kD | 16.93 | down | 0 |
| U20938 at | U20938 | 3324 | dihydropyrimidine dehydrogenase | 3.92 | down | 0.00053 |
| U21931 at | U21931 | 3326 | fructose-bisphosphatase 1 | 3.4 | down | 0.00709 |
| 1 | | | cytochrome P450, subfamily IIA (phenobarbital- | | | |
| U22029 f at | U22029 | 3327 | inducible), polypeptide 7 | 71.98 | down | 0 |
| U22961_s_at | U22961 | 3330 | albumin | 22.22 | down | 0.01531 |
|]] | | | aldehyde dehydrogenase 4 (glutamate gamma- | | | |
| U24266_at | U24266 | 3331 | semialdehyde dehydrogenase; pyrroline-5- | 14.54 | down | 0 |
| U25182_at | U25182 | 3333 | thioredoxin peroxidase (antioxidant enzyme) | 3.13 | down | 0.02543 |
| U26173_s_at | U26173 | 3335 | nuclear factor, interleukin 3 regulated | 6.4 | down | 0.00001 |
| U27460_at | U27460 | 3339 | UDP-glucose pyrophosphorylase 2 | 11.46 | down | 0.00088 |
| | | | solute carrier family 6 (neurotransmitter | | | |
| U27699_at | U27699 | 3340 | transporter, betaine/GABA), member 12 | 3.65 | down | 0.00381 |
| U28833 at | U28833 | 3341 | Down syndrome candidate region 1 | 3.3 | down | 0.00306 |
| U29953_rna1_at | U29953 | 3342 | pigment epithelium-derived factor | 15.33 | down | 0.00212 |
| U31449_at | U31449 | 3345 | transmembrane 4 superfamily member 4 | 11.53 | down | 0.00187 |
| U32576 rna1 at | U32576 | 3347 | apolipoprotein C-IV | 18.59 | down | 0.00005 |
| U32576_ma1_at | U32576 | 3347 | apolipoprotein C-IV | 5.81 | down | 0.04343 |
| U32989_at | U32989 | 3348 | tryptophan 2,3-dioxygenase | 10.07 | down | 0.02825 |
| | | | aldehyde dehydrogenase 9 (gamma- | | | |
| U34252_at | U34252 | 3350 | aminobutyraldehyde dehydrogenase, E3 | 4.31 | down | 0.00613 |
| U36922_at | U36922 | 3352 | EST | 3.06 | down | 0.01388 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|---------------------|--------|---|-------------|-----------|---------|
| | | | macrophage stimulating 1 (hepatocyte growth | | | |
| U37055_rna1_s_at | U37055 | 3353 | factor-like),macrophage stimulating, pseudogene | 18.28 | down | 0 |
| U44111_at | U44111 | 3363 | histamine N-methyltransferase | 3.5 | down | 0.00942 |
| U46499 at | U46499 | 3365 | microsomal glutathione S-transferase 1 | 4.27 | down | 0.00244 |
| U46689 at | U46689 | 3366 | aldehyde dehydrogenase 10 (fatty aldehyde | 3.5 | down | 0.01292 |
| U48296_at | U48296 | 3369 | protein tyrosine phosphatase type IVA, member 1 | 4.25 | down | 0.02037 |
| U48707_at | U48707 | 3371 | protein phosphatase 1, regulatory (inhibitor) | 14.22 | down | 0 |
| U48959 at | U48959 | 3372 | myosin, light polypeptide kinase | 3.61 | down | 0.00084 |
| U49082_at | U49082 | 3373 | transporter protein | 9.17 | down | 0.00088 |
| U49248_at | U49248 | 3374 | ATP-binding cassette, sub-family C (CFTR/MRP), | 2.06 | down | 0.00067 |
| U49352_at | U49352 | 3375 | 2,4-dienoyl CoA reductase 1, mitochondrial | 5.41 | down | 0.02371 |
| U49785_at | U49785 | 3376 | D-dopachrome tautomerase | 3.84 | down | 0.00044 |
| U50196_at | U50196 | 3377 | adenosine kinase | 3.82 | down | 0.01425 |
| U50527_s_at | U50527 | 3378 | EST | 3.93 | down | 0.00123 |
| U50929_at | U50929 | 3380 | betaine-homocysteine methyltransferase | 36.91 | down | 0 |
| U51010_s_at | U51010 | 3381 | nicotinamide N-methyltransferase | 27.79 | down | 0.00366 |
| U51333_s_at | U51333 | 3383 | hexokinase 3 (white cell) | 4.88 | down | 0.00055 |
| U51903_at | U51903 | 3387 | IQ motif containing GTPase activating protein 2 | 3.23 | down | 0.00242 |
| U53003_at | U53003 | 3388 | ES1 (zebrafish) protein, human homolog of | 4.84 | down | 0.00077 |
| U56814_at | U56814 | 3393 | deoxyribonuclease I-like 3 | 29.43 | down | 0.00003 |
| U56814_at | U56814 | 3393 | deoxyribonuclease I-like 3 | 11.36 | down | 0.00001 |
| U57721_at | U57721 | 3395 | kynureninase (L-kynurenine hydrolase) | 4.34 | down | 0.0197 |
| U60205_at | U60205 | 3401 | sterol-C4-methyl oxidase-like | 5.16 | down | 0.00061 |
| U62389_at | U62389 | 3402 | isocitrate dehydrogenase 1 (NADP+), soluble | 3.15 | down | 0.03949 |
| U66674_at | U66674 | 3408 | ATP-binding cassette, sub-family C (CFTR/MRP), | 6.13 | down | 0.00127 |
| U67963_at | N67963 | 3410 | lysophospholipase-like | 3.19 | down | 0.00029 |
| U68233_at | U68233 | 3412 | nuclear receptor subfamily 1, group H, member 4 | 9.76 | down | 0.00022 |
| U68494_at | U68494 | 3413 | EST | . 4.82 | down | 0.00881 |
| U69141_at | U69141 | 3414 | glutaryl-Coenzyrne A dehydrogenase | 3.06 | down | 0.00053 |
| U70732_rna1_at | U70732 | 3415 | glutamic-pyruvate transaminase (alanine | 27.63 | down | 0 |
| U70732_ma1_at | U70 7 32 | 3415 | glutamic-pyruvate transaminase (alanine | 5.64 | down | 0.00146 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Sea ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| U73514 at | U73514 | 3419 | hydroxyacyl-Coenzyme A dehydrogenase, type II | 3.79 | down | 0.01392 |
| U73682_at | U73682 | 3420 | meningioma expressed antigen 6 (coiled-coil | 3.08 | down | 0.01249 |
| U76376_at | U76376 | 3425 | harakiri, BCL2-interacting protein (contains only | 11.77 | down | 0.00002 |
| U77396_at | U77396 | 3426 | LPS-induced TNF-alpha factor | 4.47 | down | 0.00108 |
| U77594_at | U77594 | 3428 | retinoic acid receptor responder (tazarotene | 11.95 | down | 0.00527 |
| U78190_rna1_at | U78190 | 3431 | GTP cyclohydrolase I feedback regulatory protein | 3.94 | down | 0.034 |
| U79294 at | U79294 | 3435 | Phosphatidic acid phosphatase type 2b | 3.13 | down | 0.00129 |
| U79303_at | U79303 | 3436 | protein predicted by clone 23882 | 4.45 | down | 0 |
| U79716_at | U79716 | 3437 | reelin | 3.51 | down | 0.00053 |
| I | | | solute carrier family 9 (sodium/hydrogen | | | |
| U82108 s at | U82108 | 3441 | exchanger), isoform 3 regulatory factor 2 | 3.19 | down | 0.01545 |
| U82468 at | U82468 | 3442 | tubby like protein 1 | 3.34 | down | 0.0097 |
| U84569_at | U84569 | 3445 | chromosome 21 open reading frame 2 | 3.46 | down | 0.01844 |
| U86529_at | U86529 | 3451 | glutathione S-transferase zeta 1 | 3.41 | down | 0.0118 |
| U90544_at | U90544 | 3454 | solute carrier family 17 (sodium phosphate), | 6.42 | down | 0.00023 |
| U90545_at | U90545 | 3455 | solute carrier family 17 (sodium phosphate), | 4.14 | down | 0.00005 |
| U95090_at | 095090 | 3464 | nephrosis 1, congenital, Finnish type (nephrin) | 14.01 | down | 0.00018 |
| U95090_at | 06056N | 3464 | nephrosis 1, congenital, Finnish type (nephrin) | 7.44 | down | 0.00004 |
| W03796_at | W03796 | 3468 | EST | 8.3 | down | 0.0032 |
| W07723_at | W07723 | 3471 | EST | 3.51 | down | 0.00026 |
| rc_W15528_at | W15528 | 3475 | EST | 3.56 | down | 0.00424 |
| W20094 at | W20094 | 3477 | DKFZP586A0522 protein | 38.09 | down | 0.00188 |
| W20276 f at | W20276 | 3478 | EST | 3.82 | down | 0.00033 |
| rc W20467 f at | W20467 | 3480 | EST | 7.76 | down | 0.00008 |
| W26769 at | W26769 | 3483 | CGI-86 protein | 8.14 | down | 0.00204 |
| W26996_at | W26996 | 3484 | EST | 6.42 | down | 0.00005 |
| W27023 at | W27023 | 3485 | neuroendocrine-specific protein C like (foocen) | 6.79 | down | 0.00805 |
| W28798 at | W28798 | 3492 | phosphodiesterase 6A, cGMP-specific, rod, alpha | 7.01 | down | 0.00017 |
| W28824 at | W28824 | 3493 | EST | 15.8 | down | 0.00006 |
| W28944_at | W28944 | 3494 | EST | 37.07 | down | 0.00205 |
| rc_W31478_s_at | W31478 | 3496 | EST | 3.36 | down | 0.01511 |
| | | | | | | |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affv ID | Genbank | SealD | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|-------|--|-------------|-----------|---------|
| rc W33167 at | W33167 | 3499 | EST | 8.01 | down | 0.00026 |
| W35309 at | W35309 | 3500 | EST | 3.11 | down | 0.029 |
| W36290 s at | W36290 | 3501 | Kreisler (mouse) maf-related leucine zipper | 3.94 | down | 0.02718 |
| rc_W37382_at | W37382 | 3502 | EST | 4.37 | down | 0.00677 |
| rc_W38407_at | W38407 | 3506 | EST | 4.17 | down | 0.00392 |
| rc_W42483_at | W42483 | 3510 | EST | 3.04 | down | 0.02518 |
| rc_W42789_at | W42789 | 3515 | EST | 9.38 | down | 0.00059 |
| rc_W42996_at | W42996 | 3517 | EST | 6.18 | down | 0.00587 |
| rc_W44745_at | W44745 | 3520 | EST | 9.02 | down | 0.00276 |
| rc W45051 at | W45051 | 3521 | EST | 4.68 | down | 0.00433 |
| W45259 at | W45259 | 3522 | EST | 3.78 | down | 0.00069 |
| rc W45560 at | W45560 | 3525 | EST | 2.06 | down | 0.00996 |
| rc_W46391_at | W46391 | 3527 | alpha integrin binding protein 63 | 4.03 | down | 0.01363 |
| rc_W46404_at | W46404 | 3528 | EST | 3.73 | down | 0.00116 |
| rc_W47175_at | W47175 | 3534 | 3-prime-phosphoadenosine 5-prime- | 5.54 | down | 0.00914 |
| rc_W51951_s_at | W51951 | 3542 | dCMP deaminase | 3.14 | down | 0.01691 |
|]] | | | CD36 antigen (collagen type I receptor, | | | |
| W52581_at | W52581 | 3543 | thrombospondin receptor)-like 2 (lysosomal | 3.08 | down | 0.00091 |
| W52821_at | W52821 | 3544 | leucine aminopeptidase | 4.02 | down | 0.03787 |
| W55903_at | W55903 | 3546 | adipose differentiation-related protein; adipophilin | 9.12 | down | 0.00459 |
| rc W57821 at | W57821 | 3548 | EST | 5.03 | down | 0.00277 |
| W58540 at | W58540 | 3553 | KIAA1131 protein | 4.28 | down | 0.0184 |
| rc_W58756 at | W58756 | 3554 | EST | 12.17 | down | 0.00087 |
| rc_W60002 s_at | W60002 | 3555 | plastin 3 (T isoform) | 4.88 | down | 0.01694 |
| rc W60186 at | W60186 | 3557 | EST | 3.24 | down | 0.00228 |
| rc_W61000_at | W61000 | 3560 | EST | 3.15 | down | 0.03143 |
| rc_W61319_at | W61319 | 3561 | EST | 3.02 | down | 0.00031 |
| W61377_at | W61377 | 3562 | EST | 9.33 | down | 0.0012 |
| rc_W61378_s_at | W61378 | 3563 | EST | 8.31 | down | 0 |
| rc W63728 at | W63728 | 3565 | EST | 5.03 | down | 0.00311 |
| rc_W63785_at | W63785 | 3567 | EST | 3.91 | down | 0.04089 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affv ID | Genbank | Seq ID | Known Gene Name | Fold Change Direction | Direction | Pvalue |
|----------------|---------|--------|--|-----------------------|-----------|---------|
| rc W67147 at | W67147 | 3568 | deleted in liver cancer 1 | 9.74 | down | 0.00002 |
| rc_W67199_at | W67199 | 3569 | EST | 3.09 | down | 0.01528 |
| rc W67564 s at | W67564 | 3571 | nuclear receptor subfamily 0, group B, member 2 | 30.34 | down | 0 |
| rc W68721 f at | W68721 | 3572 | cleft lip and palate associated transmembrane | 6.34 | down | 0.01504 |
| rc W69675 at | W69675 | 3575 | EST | 4.42 | down | 0.00019 |
| rc W70115 s at | W70115 | 3576 | histidine ammonia-lyase | 11.65 | down | 0.0003 |
| rc W70313 at | W70313 | 3578 | EST | က | down | 0.00643 |
| rc W72044 at | W72044 | 3580 | insulin induced gene 1 | 24.58 | down | 0.00001 |
| rc W72079 at | W72079 | 3581 | EST | 6.36 | down | 0.00641 |
| rc_W72972_at | W72972 | 3587 | EST | 5.61 | down | 0.00939 |
| rc W73194 at | W73194 | 3590 | dermatopontin | 3.45 | down | 0.02211 |
| rc W73601 at | W73601 | 3592 | EST | 32.25 | down | 0 |
| rc W73818 at | W73818 | 3593 | EST | 4.32 | down | 0.00385 |
| rc W73889 s at | W73889 | 3594 | tetranectin (plasminogen-binding protein) | 7.45 | down | 0.00091 |
| | W74158 | 3596 | EST | 4.81 | down | 0.00233 |
| W79046 at | W79046 | 3602 | peroxisomal D3,D2-enoyl-CoA isomerase | 27.9 | down | 0.00023 |
| rc W79422 s at | W79422 | 3604 | fumarylacetoacetate | 14.94 | down | 0.00059 |
| rc W80609 at | W80609 | 3606 | EST | 4.68 | down | 0.01729 |
| W81053 at | W81053 | 3610 | EST | 4.91 | down | 0.00164 |
| rc W81079 at | W81079 | 3611 | EST | 3.31 | down | 0.0167 |
| 1 | | | protein kinase, interferon-inducible double | | | |
| W81268 at | W81268 | 3612 | stranded RNA dependent | 4.2 | down | 0.00007 |
| rc W85765 at | W85765 | 3618 | EST | 3.83 | down | 0.00379 |
| rc_W85847_at | W85847 | 3619 | EST | 7.28 | down | 0.00024 |
| rc_W85886_at | W85886 | 3621 | EST | 3.46 | down | 0.00814 |
| rc_W85888_at | W85888 | 3622 | EST | 3.99 | down | 0.00697 |
| rc_W86075_at | W86075 | 3624 | EST | 13.83 | down | 0.00057 |
| rc W86375 s at | W86375 | 3626 | EST | 19.29 | down | 0.0061 |
| rc_W86431_s at | W86431 | 3627 | protein C inhibitor (plasminogen activator inhibitor | 14.94 | down | 0.00152 |
| rc W86600 at | W86600 | 3628 | EST | 32.14 | down | 0 |
| rc_W86756_at | W86756 | 3630 | retinoid X receptor, alpha | 3.77 | down | 0.02472 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| | | | | | | 2010 |
|-------------------------|--------|------|--|-------|------|---------|
| | | | solute carrier family 25 (mitochondrial carrier; | | | |
| rc_W86850 f_at | W86850 | 3631 | citrate transporter), member 1 | 4.9 | down | 0.03105 |
| rc W87454 at | W87454 | 3632 | homogentisate 1,2-dioxygenase (homogentisate | 7.93 | down | 0.00149 |
| rc_W87480_at | W87480 | 3633 | STAT induced STAT inhibitor-2 | 3.24 | down | 0.01063 |
| rc_W87532_at | W87532 | 3634 | putative glycine-N-acyltransferase | 12.34 | down | 0.00014 |
| rc <u>W</u> 87606 s at | W87606 | 3635 | protein Z, vitamin K-dependent plasma | 9.23 | down | 0.00085 |
| rc W87781 at | W87781 | 3636 | EST | 6.1 | down | 0.00045 |
| rc_W87824_at | W87824 | 3637 | EST | 3.67 | down | 0.01559 |
| rc W88568 at | W88568 | 3638 | glycogenin 2 | 4.16 | down | 0.00111 |
| rc_W88946_at | W88946 | 3639 | putative glycine-N-acyltransferase | 58.26 | down | 0 |
| rc_W88985_at | W88985 | 3640 | KIAA0903 protein | 3.94 | down | 0.00295 |
| rc_W89178_at | W89178 | 3641 | transferrin receptor 2 | 10.85 | down | 0.00116 |
| rc W90128 s at | W90128 | 3643 | X-box binding protein 1 | 10.04 | down | 0.00018 |
| rc_W90455 s. at | W90455 | 3646 | alpha-2-macroglobulin | 10.35 | down | 0.00063 |
| | W90560 | 3647 | EST | 3.35 | down | 0.00388 |
| rc W90583 r at | W90583 | 3648 | EST | 4.35 | down | 0.00318 |
| rc_W90766_at | W90766 | 3649 | EST | 7.58 | down | 0.00057 |
| rc_W92148_s_at | W92148 | 3650 | kininogen | 51.09 | down | 0.00376 |
| rc_W92713_at | W92713 | 3654 | EST | 16.55 | down | 0.00097 |
| rc W92771 s at | W92771 | 3655 | glycine cleavage system protein H (aminomethy) | 4.36 | down | 0.0064 |
| rc_W94427_at | W94427 | 3659 | EST | 3.86 | down | 0.02649 |
| rc W94942 i at | W94942 | 3661 | dual specificity phosphatase 10 | 7.23 | down | 0.00137 |
| rc_W95041_at | W95041 | 3662 | EST | 12.79 | down | 0.00001 |
| W95795 at | W95795 | 3665 | EST | 7.8 | down | 0.00359 |
| X00129_at | X00129 | 3667 | retinol-binding protein 4, interstitial | 18.57 | down | 0.02378 |
| AFFX-HSAC07/X00351 M st | X00351 | 3668 | actin, beta | 4.76 | down | 0.0003 |
| X01038 rna1 s at | X01038 | 3669 | apolipoprotein A-I,apolipoprotein C-III | 19.97 | down | 0.0275 |
| = | X01388 | 3670 | apolipoprotein C-III | 10.9 | down | 0.01246 |
| X02160_at | X02160 | 3671 | insulin receptor | 5.29 | down | 0.0001 |
| X02160_at | X02160 | 3671 | insulin receptor | 3.27 | down | 0.0007 |
| X02176_s_at | X02176 | 3672 | complement component 9 | 9.61 | down | 0.00004 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change Direction | Direction | Pvalue |
|----------------|---------|--------|--|-----------------------|-----------|---------|
| X02750 at | X02750 | 3673 | protein C (inactivator of coagulation factors Va | 7.04 | down | 0.00079 |
| ! | | | vitronectin (serum spreading factor, somatomedin | | | |
| X03168 at | X03168 | 3674 | B, complement S-protein) | 25.88 | down | 0.00911 |
| X03350_at | X03350 | 3676 | alcohol dehydrogenase 2 (class I), beta | 19.32 | down | 0.00001 |
| X04085 rna1 at | X04085 | 3678 | catalase | 11.27 | down | 0.0002 |
| l I | ٠ | | gap junction protein, beta 1, 32kD (connexin 32, | | | |
| X04325 at | X04325 | 3679 | Charcot-Marie-Tooth neuropathy, X-linked) | 10.17 | down | 0.00061 |
| X04729 s at | X04729 | 3682 | plasminogen activator inhibitor, type I | 3.15 | down | 0.01337 |
| | | | guanine nucleotide binding protein (G protein), | | | |
| X04828 at | X04828 | 3683 | alpha inhibiting activity polypeptide 2 | 5.22 | down | 0.01278 |
| X05409 at | X05409 | 3684 | aldehyde dehydrogenase 2, mitochondrial | 3.99 | down | 0.01029 |
| X06562_at | X06562 | 3686 | growth hormone receptor | 10.87 | down | 0.00001 |
| X07173_at | X07173 | 3690 | inter-alpha (globulin) inhibitor, H2 polypeptide | 30.3 | · uwop | 0.00016 |
| | | | cytochrome P450, subfamily IID (debrisoquine, | | | |
| X07618_s_at | X07618 | 3691 | sparteine, etc., -metabolising), polypeptide 7a | 35.79 | down | 0.00065 |
| i | | | cytochrome P450, subfamily IID (debrisoquine, | | | |
| X07618 s at | X07618 | 3691 | sparteine, etc., -metabolising), polypeptide 7a | 13.58 | down | 0 |
| ! | | | cytochrome P450, subfamily IID (debrisoquine, | | | |
| X07619 s at | X07619 | 3692 | sparteine, etc., -metabolising), polypeptide 7a | 4.96 | down | 0.00004 |
| X07732 at | X07732 | 3693 | hepsin (transmembrane protease, serine 1) | 28.21 | down | 0 |
| X07767_at | X07767 | 3694 | protein kinase, cAMP-dependent, catalytic, alpha | 5.94 | down | 0.00028 |
| X12662 ma1 at | X12662 | 3697 | arginase, liver | 20.59 | down | 0 |
| X13227 at | X13227 | 3698 | D-amino-acid oxidase | 4.81 | down | 0.0002 |
| X13334_at | X13334 | 3699 | CD14 antigen | 12.77 | down | 0.00887 |
| | | | cytochrome P450, subfamily IIA (phenobarbital- | | | |
| X13930 f at | X13930 | 3700 | inducible), polypeptide 6 | 38.52 | down | 0 |
| X14690 s at | X14690 | 3703 | pre-alpha (globulin) inhibitor, H3 polypeptide | 16.07 | down | 0.00438 |
| X14787 at | X14787 | 3704 | thrombospondin 1 | 3.69 | down | 0.01115 |
| I | | | acetyl-Coenzyme A acyltransferase 1 | | | |
| X14813_at | X14813 | 3705 | (peroxisomal 3-oxoacyl-Coenzyme A thiolase) | 11.61 | down | 0.0002 |
| X15422_at | X15422 | 3708 | mannose-binding lectin (protein C) 2, soluble | 8.29 | down | 0.00015 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| ATTY ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|---------|--------|--|-------------|-----------|---------|
| X16260_s_at | X16260 | 3710 | inter-alpha (globulin) inhibitor, H1 polypeptide | 44.65 | down | 0.00933 |
| X16260_s_at | X16260 | 3710 | inter-alpha (globulin) inhibitor, H1 polypeptide | 26.39 | down | 0.0004 |
| X16349_s_at | X16349 | 3712 | sex hormone-binding globulin | 9.34 | down | 0.00007 |
| X17025_at | X17025 | 3715 | isopentenyl-diphosphate delta isomerase | 3.13 | down | 0.0043 |
| | | | paired basic amino acid cleaving enzyme (furin, | | | |
| X17094_at | X17094 | 3717 | membrane associated receptor protein) | 8.5 | down | 0 |
| X52150 rna1 s at | X52150 | 3722 | arylsulfatase A | 3.51 | down | 0.00113 |
| X52520 at | X52520 | 3723 | tyrosine aminotransferase | 13.51 | down | 0.00002 |
| X52541_at | X52541 | 3724 | early growth response 1 | 3.77 | down | 0.00894 |
| l | | | alanine-glyoxylate aminotransferase (oxalosis I; | | | |
| X53414 at | X53414 | 3728 | hyperoxaluria I; glycolicaciduria; serine-pyruvate | 13.87 | down | 0.00003 |
| X53595 s at | X53595 | 3729 | apolipoprotein H (beta-2-glycoprotein I) | 27.28 | down | 0.0066 |
| X54380 at | X54380 | 3730 | pregnancy-zone protein | 8.44 | down | 0.00059 |
| X55283 rna1_s_at | X55283 | 3734 | asialoglycoprotein receptor 2 | 84.99 | down | 0.00084 |
| X56411_ma1_at | X56411 | 3737 | alcohol dehydrogenase 4 (class II), pi polypeptide | | down | 0.00144 |
| X56692_at | X56692 | 3739 | C-reactive protein, pentraxin-related | 15.15 | down | 0.01884 |
| X57025_at | X57025 | 3742 | insulin-like growth factor 1 (somatomedin C) | 4.72 | down | 0.00087 |
| X58022_at | X58022 | 3747 | corticotropin releasing hormone-binding protein | 4.09 | down | 0.00076 |
| X58528_s_at | X58528 | 3748 | ATP-binding cassette, sub-family D (ALD), | 4.91 | down | 0.00728 |
| X59766_at | X29766 | 3749 | alpha-2-glycoprotein 1, zinc | 12.96 | down | 0.00043 |
| • | | | cytochrome P450, subfamily XXVIIA (steroid 27- | | | |
| X59812_at | X59812 | 3750 | hydroxylase, cerebrotendinous xanthomatosis), | 4.65 | down | 0.036 |
| X59834_at | X59834 | 3751 | glutamate-ammonia ligase (glutamine synthase) | 4.29 | down | 0.00255 |
| X60673_s_at | X60673 | 3752 | adenylate kinase 3 | 8.3 | down | 0.00016 |
| X61123_at | X61123 | 3753 | B-cell translocation gene 1, anti-proliferative | 3.2 | down | 0.00796 |
| X62822_at | X62822 | 3758 | sialyltransferase 1 (beta-galactoside alpha-2,6- | 4.89 | down | 0.00274 |
| X63359_at | X63359 | 3759 | UDP glycosyltransferase 2 family, polypeptide | 13.66 | down | 0.00051 |
| X64877_at | X64877 | 3766 | H factor (complement)-like 3 | 14.6 | down | 0 |
| X64877_s_at | X64877 | 3766 | H factor (complement)-like 3 | 12.33 | down | 0 |
| X65727_cds2_s_at | X65727 | 3768 | glutathione S-transferase A2, glutathione S- | 73.64 | down | 0 |
| X65962 s at | X65962 | 3769 | cytochrome P450, subfamily IIC (mephenytoin 4- | 5.47 | down | 0.00014 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|-------------------|---------|--------|---|-------------|-----------|----------|
| X67235_s_at | X67235 | 3773 | hematopoietically expressed homeobox | 3.17 | down | 0.0045 |
| X67491_f_at | X67491 | 3776 | glutamate dehydrogenase 1 | 7.45 | down | 0.00019 |
| X68277_at | X68277 | 3777 | dual specificity phosphatase 1 | 6.68 | down | 0.0036 |
| X68679_s_at | X68679 | 3779 | complement factor H related 3, complement factor | 38.22 | down | 0.00036 |
| X68733_rna1_at | X68733 | 3781 | alpha-1-antichymotrypsin | 7.87 | down | 0.03266 |
| X72012_at | X72012 | 3789 | endoglin (Osler-Rendu-Weber syndrome 1) | 4.2 | down | 0.005 |
| X72177_rna1_at | X72177 | 3790 | complement component 6 | 12.94 | down | 0.00011 |
| X75252_at | X75252 | 3793 | prostatic binding protein | 3.78 | down | 0.00231 |
| X76105_at | X76105 | 3794 | death-associated protein | 3.02 | down | 0.00944 |
| X76648_at | X76648 | 3796 | glutaredoxin (thioltransferase) | 4 | down | 0.00211 |
| X76717_at | X76717 | 3797 | metallothionein 1L | 8.09 | down | 0.00025 |
| X77548_at | X77548 | 3798 | nuclear receptor coactivator 4 | 3.76 | down | 0.00758 |
| X78706_at | X78706 | 3801 | carnitine acetyltransferase | 4.65 | down | 0.00442 |
| X78992_at | X78992 | 3802 | butyrate response factor 2 (EGF-response factor | 4.69 | down | 0.01995 |
| | | | prion protein (p27-30) (Creutzfeld-Jakob disease, | | | |
| X83416_s_at | X83416 | 3811 | Gerstmann-Strausler-Scheinker syndrome, fatal | 3.28 | down | 0.00221 |
| | | | 3-hydroxy-3-methylglutaryl-Coenzyme A synthase | | | |
| X83618_at | X83618 | 3813 | 2 (mitochondrial) | 6.27 | down | 0.02099 |
| X85116_rna1_s_at | X85116 | 3814 | erythrocyte membrane protein band 7.2 | 6.37 | down | 0.00356 |
| X86401_s_at | X86401 | 3815 | glycine amidinotransferase (L-arginine:glycine | 22.7 | down | 0 |
| X87344_cds10_r_at | X87344 | 3817 | EST | 3.04 | down | 0.02779 |
| X90579_s_at | X90579 | 3819 | EST | 29.82 | down | 0.00273 |
| X90999_at | 66606X | 3820 | hydroxyacyl glutathione hydrolase; glyoxalase 2 | 5.27 | down | 0.00047 |
| X91148_at | X91148 | 3821 | microsomal triglyceride transfer protein (large | 5.91 | down | 0.00045 |
| X92720_at | X92720 | 3826 | phosphoenolpyruvate carboxykinase 2 | 7.6 | down | 0.00001 |
| X92744_at | X92744 | 3827 | defensin, beta 1 | 4.96 | down | 0.01804 |
| X94563_xpt2_r_at | X94563 | 3831 | EST | 3.52 | down | 0.00928. |
| X95190_at | X95190 | 3832 | acyl-Coenzyme A oxidase 2, branched chain | 12.81 | down | 0 |
| X95384_at | X95384 | 3833 | translational inhibitor protein p14.5 | 17.69 | down | 0.00048 |
| X95715_at | X95715 | 3835 | ATP-binding cassette, sub-family C (CFTR/MRP), | 6.78 | down | 0.00045 |
| X95876_at | X95876 | 3836 | G protein-coupled receptor 9 | 3.12 | down | 0.001 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name F | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|-----------|---------|
| X96752 at | X96752 | 3837 | L-3-hydroxyacyl-Coenzyme A dehydrogenase, | 4.94 | down | 0.00359 |
| X97324 at | X97324 | 3839 | adipose differentiation-related protein; adipophilin | 5.97 | down | 0.04638 |
| X98337_s_at | X98337 | 3840 | complement factor H related 3, complement factor | 13.5 | down | 0.00001 |
| Y00097_s_at | Y00097 | 3844 | annexin A6 | 3.4 | down | 0.00233 |
| Y00317 at | Y00317 | 3845 | UDP glycosyltransferase 2 family, polypeptide B4 | 18.34 | down | 0.00001 |
| Y00317_at | Y00317 | 3845 | UDP glycosyltransferase 2 family, polypeptide B4 | 8.97 | down | 0.00025 |
| Y00318_at | Y00318 | 3846 | I factor (complement) | 10 | down | 0.00019 |
| Y00339 s at | Y00339 | 3847 | carbonic anhydrase II | 6.89 | down | 0 |
| Y00451_s_at | Y00451 | 3848 | aminolevulinate, delta-, synthase 1 | 10.52 | down | 0.00107 |
| Y08374 rna1 at | Y08374 | 3853 | chitinase 3-like 1 (cartilage glycoprotein-39) | 3.5 | down | 0.04208 |
| Y08409 at | Y08409 | 3854 | thyroid hormone responsive SPOT14 (rat) | 5.84 | down | 0.00455 |
| Y09616_at | Y09616 | 3857 | carboxylesterase 2 (intestine, liver) | 18.78 | down | 0.00026 |
| Y10032_at | Y10032 | 3858 | serum/glucocorticoid regulated kinase | 4.24 | down | 0.00148 |
| Y10659_at | Y10659 | 3859 | interleukin 13 receptor, alpha 1 | 4.22 | down | 0.00061 |
| Y10659 at | Y10659 | 3859 | interleukin 13 receptor, alpha 1 | 3.17 | down | 0.00095 |
| Y12711_at | Y12711 | 3861 | progesterone binding protein | 14.83 | down | 0.00285 |
| Z11559_at | Z11559 | 3862 | iron-responsive element binding protein 1 | 4.3 | down | 0.00066 |
| Z11737_at | Z11737 | 3863 | flavin containing monooxygenase 4 | 3.84 | down | 0.00043 |
| Z11737 at | Z11737 | 3863 | flavin containing monooxygenase 4 | 3.67 | down | 0.00632 |
| Z11793_at | Z11793 | 3864 | selenoprotein P, plasma, 1 | 9.94 | down | 0.00021 |
| l | | | branched chain keto acid dehydrogenase E1, | | | |
| Z14093_at | Z14093 | 3865 | alpha polypeptide (maple syrup urine disease) | 3.24 | down | 0.00301 |
| Z20777_at | Z20777 | 3866 | EST | 29.59 | down | 0.00044 |
| Z24459 rna1 at | Z24459 | 3869 | mature T-cell proliferation 1 | 5.29 | down | 0.00001 |
| Z24725 at | Z24725 | 3870 | mitogen inducible 2 | 7.9 | down | 0 |
| Z26491 s at | Z26491 | 3873 | catechol-O-methyltransferase | 3.08 | down | 0.00877 |
| ! ! | | | aldo-keto reductase family 1, member D1 (delta 4. | | | |
| Z28339_at | Z28339 | 3875 | 3-ketosteroid-5-beta-reductase) | 24.66 | down | 0 |
| Z29481_at | Z29481 | 3877 | 3-hydroxyanthranilate 3,4-dioxygenase | 6.39 | down | 0.00029 |
| Z29481_at | Z29481 | 3877 | 3-hydroxyanthranilate 3,4-dioxygenase | 3.64 | down | 0.00096 |
| Z30425_at | Z30425 | 3878 | nuclear receptor subfamily 1, group I, member 3 | 26.64 | down | 0 |

Table 6B. Down regulated in metastatic cancers versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|---------|--------|---|-------------|-----------|---------|
| Z30425_at | Z30425 | 3878 | nuclear receptor subfamily 1, group I, member 3 | 7.88 | down | 0.00006 |
| Z31357_at | Z31357 | 3880 | cysteine dioxygenase, type I | 11.2 | down | 0.0001 |
| Z31690_s_at | Z31690 | 3881 | lipase A, lysosomal acid, cholesterol esterase | 3.83 | down | 0.00103 |
| rc_Z38161_at | Z38161 | 3884 | EST | 4.38 | down | 0.0011 |
| rc Z38192 s at | Z38192 | 3885 | EST | 3.35 | down | 0.00184 |
| rc_Z38435_at | Z38435 | 3890 | ribosomal protein L21 | 3.12 | down | 0.03617 |
| rc Z38777 f at | Z38777 | 3895 | nuclear receptor binding factor-2 | 3.28 | down | 0.022 |
| rc_Z39059_at | Z39059 | 3899 | EST | 5.19 | down | 0.0014 |
| rc_Z39406_at | Z39406 | 3905 | nuclear receptor co-repressor 1 | 4.18 | down | 0.00439 |
| rc_Z39431_at | Z39431 | 3907 | KIAA1086 protein | 3.68 | down | 0.0013 |
| rc Z39476 at | Z39476 | 3908 | EST | 5.9 | down | 0.00687 |
| rc_z39622_s_at | Z39622 | 3910 | EST | 4.4 | down | 0.00001 |
| rc Z39818 at | Z39818 | 3912 | EST | 3.26 | down | 0.00089 |
| rc_Z39833_at | Z39833 | 3913 | GTP-binding protein | 16.89 | down | 0.00034 |
| rc_Z39976_at | Z39976 | 3915 | EST | 5.76 | down | 0.00012 |
| rc_Z39978_at | Z39978 | 3916 | EST | 3.56 | down | 0.04051 |
| Z40192 | Z40192 | 3918 | EST | 3.89 | down | 0.00223 |
| rc_Z40259_s_at | Z40259 | 3919 | EST | 8.18 | down | 0.00002 |
| rc_Z40305_at | Z40305 | 3920 | EST | 6.45 | down | 0.00001 |
| rc_Z40715_at | Z40715 | 3923 | delta-6 fatty acid desaturase | 18.68 | down | 0.0007 |
| rc_Z40902_at | Z40902 | 3926 | SEC14 (S. cerevisiae)-like 2 | 12.87 | down | 0.00001 |
| rc_Z41042_at | Z41042 | 3928 | EST | 3.63 | down | 0.00943 |
| Z47553_at | Z47553 | 3939 | flavin containing monooxygenase 5 | 6.17 | down | 0.00011 |
| Z48199_at | Z48199 | 3942 | syndecan 1 | 4.43 | down | 0.00408 |
| Z48475_at | Z48475 | 3943 | glucokinase (hexokinase 4) regulatory protein | 13.84 | down | 0 |
| Z49269_at | Z49269 | 3945 | small inducible cytokine subfamily A (Cys-Cys), | 18.46 | down | 0.00001 |
| Z49878_at | Z49878 | 3946 | guanidinoacetate N-methyltransferase | 13.96 | down | 0.00021 |
| Z69923_at | Z69923 | 3948 | HGF activator | 11.37 | down | 0.00053 |
| Z80345_rna1_s_at | Z80345 | 3951 | acyl-Coenzyme A dehydrogenase, C-2 to C-3 | 5.93 | down | 0.0024 |
| Z84718 cds1 at | Z84718 | 3952 | EST | 3.26 | down | 0.02252 |
| Z84721 cds2 at | Z84721 | 3953 | hemoglobin, zeta | 8.77 | down | 0.01446 |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Nown Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| rc_AA005358_at | AA005358 | 14 | EST | 5.51 | down | 0.00059 |
| rc_AA007395_s_at | AA007395 | 17 | alcohol dehydrogenase 4 (class II), pi polypeptide | 37.78 | down | 0.00939 |
| · rc_AA007629_at | AA007629 | 19 | EST | 5.56 | down | 0.00005 |
| rc_AA010205_at | AA010205 | 23 | EST | 5.71 | down | 0.00014 |
| rc_AA010605_s_at | AA010605 | 26 | 4-hydroxypheny/pyruvate dioxygenase | 25.52 | down | 0.00855 |
| rc_AA010619_at | AA010619 | 27 | EST | 5.28 | down | 0.002 |
| rc_AA018922_s_at | AA018922 | 40 | core promoter element binding protein | 3.39 | down | 0.01801 |
| rc_AA035245_s_at | AA035245 | 79 | aldehyde oxidase 1 | 7.97 | down | 0.02387 |
| rc_AA035457_at | AA035457 | 80 | EST | 3.41 | down | 0.00968 |
| rc_AA039335_s_at | AA039335 | 83 | coagulation factor XII (Hageman factor) | 5.33 | down | 0.03807 |
| rc_AA039616_at | AA039616 | 90 | EST | 3.84 | down | 0.00997 |
| rc_AA046457_at | AA046457 | 111 | EST | 3.05 | down | 0.02078 |
| rc_AA046747_at | AA046747 | 114 | EST | 4.77 | down | 0.00023 |
| rc_AA056482_at | AA056482 | 141 | EST | 3.06 | down | 0.01313 |
| rc_AA057678_at | AA057678 | 143 | EST | 6.63 | down | 0.00089 |
| rc_AA069696_at | AA069696 | 150 | EST | 3.07 | down | 0.01569 |
| rc_AA070191_at | AA070191 | 154 | EST | 3.24 | down | 0.00216 |
| rc_AA074885_at | AA074885 | 161 | macrophage receptor with collagenous structure | 10.88 | down | 0.00087 |
| rc_AA076326_at | AA076326 | 170 | SEC14 (S. cerevisiae)-like 2 | 3.85 | down | 0.0349 |
| rc_AA076383_at | AA076383 | 171 | EST | 4.65 | down | 0.00593 |
| rc_aa084668_at | AA084668 | 180 | ubiquitin-like 3 | 3.31 | down | 0.02055 |
| rc_AA085987_s_at | AA085987 | 183 | UDP glycosyltransferase 1 | 4.74 | down | 0.03035 |
| AA090257_at | AA090257 | 190 | superoxide dismutase 2, mitochondrial | 3.03 | down | 0.02774 |
| AA090439_at | AA090439 | 192 | ribosomal protein S6 | 5.11 | down | 0.01108 |
| rc_AA099225_at | AA099225 | 206 | EST | 6.59 | down | 0.00064 |
| rc_AA100026_at | AA100026 | 211 | EST | 4.18 | down | 0.00567 |
| rc_AA112101_f_at | AA112101 | 222 | EST | 4.36 | down | 0.03175 |
| rc_AA122345_f_at | AA122345 | 238 | glutamate dehydrogenase 1 | 3.75 | down | 0.01058 |
| rc_AA129390_at | AA129390 | 262 | EST | 4.03 | down | 0.00128 |
| rc_AA142849_at | AA142849 | 306 | EST | 3.45 | down | 0.03495 |
| rc_aa147646_s_at | AA147646 | 317 | DKFZP586A0522 protein | 3.19 | down | 0.00508 |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| 75 AAE0776 24 | Gendank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| מייסטועל זו | AA150776 | 330 | EST | 6.17 | down | 0.00004 |
| rc_AA151676_at | AA151676 | 337 | peptidyl arginine deiminase, type II | 3.85 | down | 0.00875 |
| | | | aldo-keto reductase family 7, member A2 (aflatoxin | | | |
| rc_AA157799_at | AA157799 | 348 | aldehyde reductase) | 3.08 | down | 0.00207 |
| rc_AA164586_s_at | AA164586 | 328 | estrogen receptor 1 | 3.56 | down | 0.01231 |
| rc_AA167565_at | AA167565 | 362 | EST | 3.81 | down | 0.04057 |
| rc_AA172372_at | AA172372 | 370 | EST | 5.12 | down | 0.00032 |
| rc_AA182030_at | AA182030 | 387 | EST | 3.51 | down | 0.0403 |
| AA188921_at | AA188921 | 393 | similar to Caenorhabditis elegans protein C42C1.9 | 3.38 | down | 0.00862 |
| rc_AA194997_s_at | AA194997 | 412 | EST | 4.8 | down | 0.00153 |
| rc_AA196287_at | AA196287 | 420 | EST | 4.86 | down | 0.01656 |
| rc_AA210850_at | AA210850 | 431 | EST | 3.12 | down | 0.00288 |
| rc_AA223902_at | AA223902 | 450 | EST | 4.22 | down | 0.01315 |
| rc_AA232114_s_at | AA232114 | 463 | epoxide hydrolase 2, cytoplasmic | 6.18 | down | 0.00231 |
| rc_AA233152_at | AA233152 | 467 | EST | 5.8 | down | 0.00272 |
| rc_AA233837_at | AA233837 | 474 | EST | 3.46 | down | 0.01365 |
| rc_AA235310_at | AA235310 | 496 | EST | 7.08 | down | 0.04056 |
| rc_AA236401_at | AA236401 | 510 | EST | 5.31 | down | 0.01787 |
| rc_AA236455_r_at | AA236455 | 512 | EST | 6.73 | down | 0.02418 |
| rc_AA236455_s_at | AA236455 | 512 | EST | 5.1 | down | 0.00307 |
| rc_AA253216_at | AA253216 | 561 | EST | 4.14 | down | 0.0014 |
| rc_AA253369_s_at | AA253369 | 563 | EST | 5.64 | down | 0.00478 |
| rc_AA256367_s_at | AA256367 | 579 | paraoxonase 3 | 8.37 | down | 0.02326 |
| rc_AA258350_at | AA258350 | 592 | EST | 4.1 | down | 0.02962 |
| rc_AA279676_s_at | AA279676 | 630 | deoxyribonuclease I-like 3 | 10.52 | down | 0.00181 |
| rc_AA282061_at | AA282061 | 652 | KIAA0962 protein | 3.46 | down | 0.00698 |
| rc_AA282886_at | AA282886 | 663 | EST | 3.29 | down | 0.00025 |
| rc_AA284795_at | AA284795 | 678 | phosphatidylethanolamine N-methyltransferase | 2.7 | down | 0.00004 |
| rc_AA285053_at | AA285053 | 681 | EST | ιO | down | 0.00718 |
| rc_AA287122_at | AA287122 | 989 | EST | 5.21 | down | 0.00523 |
| rc_AA287566_at | AA287566 | 069 | KIAA0187 gene product | 6.99 | down | 0.00023 |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank Seq ID | ID Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------------|---|-------------|-----------|---------|
| rc_AA291749_s_at | AA291749 70 | 703 estrogen receptor 1 | 5.06 | down | 0.00044 |
| rc_AA297532_f_at | AA297532 72 | 725 EST | 5.01 | down | 0.00745 |
| rc_AA343142_at | AA343142 751 | 31 EST | 5.79 | down | 0.02747 |
| | | fatty-acid-Coenzyme A ligase, long-chain 1, fatty-acid- | | | |
| rc_AA348922_s_at | AA348922 78 | 758 Coenzyme A ligase, long-chain 2 | 11.4 | down | 0.00848 |
| AA376875_at | AA376875 7. | 770 monoamine oxidase A | 3.45 | down | 0.00105 |
| rc_AA377087_at | AA377087 7. | 71 EST | 4.61 | dowń | 0.01616 |
| . AA397841_at | AA397841 78 | 30 EST | 3.29 | down | 0.00825 |
| rc_AA398892_at | AA398892 8(| 800 similar to yeast BET3 (S. cerevisiae) | 4.33 | down | 0.01326 |
| AA400177_at | AA400177 8(| | 3.21 | down | 0.03901 |
| rc_AA401376_at | AA401376 82 | 829 EST | 3.39 | down | 0.01403 |
| rc_AA401562_s_at | AA401562 83 | 830 EST | 7.97 | down | 0.00527 |
| rc_AA402224_at | AA402224 83 | | 3.66 | down | 0.0033 |
| rc_AA404487_at | AA404487 85.1 | | 3.01 | down | 0.0059 |
| rc_AA417046_at | AA417046 9° | 915 fatty-acid-Coenzyme A ligase, very long-chain 1 | 8.49 | down | 0.02476 |
| | | small inducible cytokine subfamily B (Cys-X-Cys), | | | |
| rc_AA426640_at | AA426640 9(| 969 member 14 (BRAK) | 4.8 | down | 0.00539 |
| rc_AA428325_at | AA428325 98 | 988 EST | 4.09 | down | 0.02486 |
| rc_AA433946_at | • | 1033 EST | 10.24 | down | 0.00663 |
| rc_AA435746_f_at | AA435746 10 | 1043 GTPase activating protein-like | 4.21 | down | 0.03192 |
| rc_AA435985_at | AA435985 10 | 1049 EST | 3.86 | down | 0.01713 |
| AA442334_at | | | 3.02 | down | 0.01936 |
| rc_AA446864_at | AA446864 10 | 1095 EST | 11.57 | down | 0.0001 |
| rc_AA448002_at | AA448002 11 | 1113 putative type II membrane protein | 14.14 | down | 0 |
| | | FXYD domain-containing ion transport regulator 1 | - | | |
| rc_AA448300_at | AA448300 11 | 1116 (phospholemman) | 9.27 | down | 0.00108 |
| rc_AA450114_at | AA450114 11 | | 3.29 | down | 0.01171 |
| rc_AA450127_at | AA450127 11 | 1132 growth arrest and DNA-damage-inducible, beta | 3.37 | down | 0.00647 |
| rc_AA453988_at | AA453988 11 | 1160 methionine adenosyltransferase I, alpha | 7.78 | down | 0.02695 |
| rc_AA454733_s_at | AA454733 11 | 1169 EST | 5.73 | down | 0.00748 |
| rc_AA455367_at | AA455367 11 | 1176 DKFZP586F1018 protein | 3.47 | down | 0.00138 |
| | | | | | |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|-----------|
| | | | butyrobetaine (gamma), 2-oxoglutarate dioxygenase | | | |
| rc AA455988 at | AA455988 | 1184 | (gamma-butyrobetaine hydroxylase) | 12.51 | down | 0 |
| rc_AA458652_at | AA458652 | 1202 | EST | 5.03 | down | 0.00065 |
| 1 | | | solute carrier family 22 (extraneuronal monoamine | | | |
| rc AA460012 at | AA460012 | 1224 | transporter), member 3 | 3.73 | down | 0.0313 |
| rc_AA460449_at | AA460449 | 1228 | EST | 3.53 | down | 0.01247 |
| rc_AA460661_at | AA460661 | 1229 | EST | 5.46 | down | 0.00151 |
| rc_AA461444_at | AA461444 | 1239 | EST | 3.19 | down | 0.02844 |
| rc AA465233 s at | AA465233 | 1269 | succinate-CoA ligase, GDP-forming, beta subunit | 3.19 | down | 0.00036 |
| rc AA477119 at | AA477119 | 1289 | EST | 4.62 | down | 0.00072 |
| AA477919 at | AA477919 | 1293 | EST | 3.9 | down | 0.00265 |
| rc AA478298 s at | AA478298 | 1297 | adipose specific 2 | 5.29 | down | 0.00943 |
| rc_AA480991 s_at | AA480991 | 1323 | EST | 4.83 | down | 0.03498 |
| AA486511 at | AA486511 | 1349 | EST | 3.9 | down | 0.01409 |
| rc AA490620 at | AA490620 | 1378 | EST | 6.25 | down | 0.03613 |
| rc_AA599472_at | AA599472 | 1451 | succinate-CoA ligase, GDP-forming, beta subunit | 3.31 | down | 0.02619 |
| rc_AA599814_at | AA599814 | 1456 | EST | 4.09 | down | 0.00235 |
| rc AA599937 s at | AA599937 | 1458 | insulin-like growth factor-binding protein 4 | 6.31 | down | 0.0477 |
| rc AA608802 at | AA608802 | 1470 | EST | 3.44 | down | 0.01709 |
| rc_AA608837_at | AA608837 | 1472 | EST | 5.15 | down | 0.00005 |
| rc_AA609519_at | AA609519 | 1482 | EST | 5.23 | down | 0.00068 |
| rc AA609537 s at | AA609537 | 1483 | hepatic leukemia factor | 4.88 | down | 0.00118 |
| rc AA621131 at | AA621131 | 1513 | EST | 4.57 | down | 0.03867 |
| rc_AA621209_at | AA621209 | 1516 | similar to Caenorhabditis elegans protein C42C1.9 | 4.01 | down | 0.00563 |
| C02386 s at | C02386 | 1561 | hypothetical protein | 3.66 | down | 0.00673 |
| C02460 at | C02460 | 1562 | EST | 3.92 | down | 0.03073 |
| rc C20653 at | C20653 | 1578 | EST | 4.32 | down | . 0.00718 |
| rc_C20810_at | C20810 | 1579 | EST | 3.57 | down | 0.02116 |
| rc_C21130_at | C21130 | 1583 | EST | 3.24 | down | 0.03355 |
| l l | | | cytochrome P450, subfamily IIIA (niphedipine oxidase), | | | |
| D00003_at | D00003 | 1586 | polypeptide 3 | 9.46 | down | 0.00001 |
| | | | | | | |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| 1 | Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue | |
|---|----------------|---------|--------|--|-------------|-----------|---------|--|
| 1 | | | | cytochrome P450, subfamily IIIA (niphedipine oxidase), | | | | |
| | D00003_s_at | D00003 | 1586 | polypeptide 3 | 6.8 | down | 0.01328 | |
| | | | | cytochrome P450, subfamily IIIA (niphedipine oxidase), | | | | |
| | | | | polypeptide 3, cytochrome P450, subfamily IIIA | | | | |
| | | | | (niphedipine oxidase), polypeptide 5, cytochrome P450, | | | | |
| | D00408_s_at | D00408 | 1589 | subfamily IIIA, polypeptide 7 | 3.58 | down | 0.02048 | |
| | D10040_at | D10040 | 1593 | fatty-acid-Coenzyme A ligase, long-chain 2 | 4.15 | down | 0.02947 | |
| | rc_D11756_f_at | D11756 | 1596 | EST | 4.08 | down | 0.02972 | |
| | | | | cytochrome P450, subfamily IVF, polypeptide | | | | |
| | | | | 2, cytochrome P450, subfamily IVF, polypeptide 3 | | | | |
| | D12620_s_at | D12620 | 1601 | (leukotriene B4 omega hydroxylase) | 6.03 | down | 0.03947 | |
| | | | | cytochrome P450, subfamily IVF, polypeptide | | | · | |
| | | | | 2, cytochrome P450, subfamily IVF, polypeptide 3 | | | | |
| | D12620_s_at | D12620 | 1601 | (leukotriene B4 omega hydroxylase) | 4.7 | down | 0.04091 | |
| | D13243_s_at | D13243 | 1602 | pyruvate kinase, liver and RBC | 5.58 | down | 0.04029 | |
| • | D13705_s_at | D13705 | 1610 | cytochrome P450, subfamily IVA, polypeptide 11 | 3.3 | down | 0.0051 | |
| | D14012_s_at | D14012 | 1612 | HGF activator | 7.27 | down | 0.00145 | |
| | D31628_s_at | D31628 | 1646 | 4-hydroxyphenylpyruvate dioxygenase | 13.18 | down | 0.02064 | |
| | rc_D45529_at | D45529 | 1662 | EST | 3.01 | down | 0.03105 | |
| | D49357_at | D49357 | 1665 | methionine adenosyltransferase I, alpha | 4.85 | down | 0.04435 | |
| | rc_D62518_at | D62518 | 1708 | EST | 5.96 | down | 0.00027 | |
| | | | | ficolin (collagen/fibrinogen domain-containing lectin) 2 | | | | |
| | D63160_at | D63160 | 1709 | (hucolin) | 3.72 | down | 0.00312 | |
| | D78011_at | D78011 | 1717 | dihydropyrimidinase | 5.54 | down | 0.0312 | |
| | D78725_at | D78725 | 1720 | KIAA0914 gene product | 3.19 | down | 0.01083 | |
| | D79276_at | D79276 | 1722 | succinate-CoA ligase, GDP-forming, beta subunit | 4.34 | down | 0.00836 | |
| • | D90042_at | D90042 | 1767 | N-acetyltransferase 2 (arylamine N-acetyltransferase) | 3.79 | down | 0.00697 | |
| • | rc_F02028_at | F02028 | 1774 | EST | 3.15 | down | 0.00902 | |
| | rc_F02245_at | F02245 | 1776 | monoamine oxidase A | 3.51 | down | 0.01692 | |
| | | | | matrix metalloproteinase 2 (gelatinase A, 72kD | | | | |
| | rc_F03969_at | F03969 | 1785 | gelatinase, 72kD type IV collagenase) | 3.36 | down | 0.01685 | |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|----------------------------------|--------|---|-------------|-----------|---------|
| rc H02855 at | H02855 | 1832 | EST | 4.29 | down | 0.0138 |
| rc H05704 s at | | 1848 | EST | 3.07 | down | 0.00363 |
| rc_H06935_s_a | at H06935 | 1855 | electron-transferring-flavoprotein dehydrogenase | 4.04 | down | 0.01498 |
| rc H08102.at | H08102 | 1858 | breast cell glutaminase | 12.85 | down | 0.0424 |
| rc_H09317_at | | 1864 | EST | 3.2 | down | 0.00914 |
| l I | | | methylenetetrahydrofolate dehydrogenase (NADP+ | | | |
| | | | dependent), methenyltetrahydrofolate cyclohydrolase, | | | |
| rc H10779 s at | at H10779 | 1872 | formyltetrahydrofolate synthetase | 3.09 | down | 0.0496 |
| rc H20543 at | H20543 | 1897 | DKFZP586B1621 protein | 13.06 | down | 0.00218 |
| rc_H27330_at | H27330 | 1909 | EST | 3.37 | down | 0.01318 |
| rc_H29568_at | H29568 | 1914 | EST | 2 | down | 0.00426 |
| rc_H55759_at | H55759 | 1949 | EST | 4.36 | down | 0.0398 |
| rc H57060 s at | _ | 1954 | EST | 7.57 | down | 0.00875 |
| rc H57816 at | | 1957 | EST | 3.09 | down | 0.01327 |
| rc_H58673_at | H58673 | 1959 | EST | 15.49 | down | 0.00002 |
| rc h58692 s at | at H58692 | 1960 | formyltetrahydrofolate dehydrogenase | 20.18 | down | 0.00485 |
| rc H59136 at | H59136 | 1962 | EST | 6.63 | down | 0.00033 |
| rc_H62212_at | H62212 | 1969 | telomeric repeat binding factor 2 | 3.23 | down | 0.00513 |
| H66367 at | H66367 | 1977 | EST | 3.84 | down | 0.00133 |
| rc H66840 at | H66840 | 1978 | EST | 3.34 | down | 0.01884 |
| rc_H77597_f_at | t H77597 | 2000 | metallothionein 1H | 9.01 | down | 0.00022 |
|] | | | ficolin (collagen/fibrinogen domain-containing) 3 (Hakata | | | |
| rc H80901 s at | at H80901 | 2002 | antigen) | 18.59 | down | 0 |
| rc_H81070_f_at | at H81070 | 2006 | RNA helicase-related protein | 39.64 | down | 0.00002 |
| rc H87765 at | H87765 | 2017 | KIAA0626 gene product | 3.94 | down | 0.00123 |
| H93246 s at | H93246 | 2035 | EST | 4.14 | down | 0.00058 |
| rc H93381 at | | 2036 | EST | 8.62 | down | 0.01271 |
| rc_H99727_at | t H99727 | 2080 | adipose differentiation-related protein; adipophilin | 3.91 | down | 0.00325 |
| HG1428-HT1428 | HG1428-HT1428 s at HG1428-HT1428 | | hemoglobin, beta | 8.98 | down | 0.02071 |
| HG2379-HT3996 | HG2379-HT3996 s at HG2379-HT3996 | | serine hydroxymethyltransferase 1 (soluble) | 3.81 | down | 0.01837 |
| HG2730-HT2827_ | HG2730-HT2827_s_at HG2730-HT2827 | | fibrinogen, A alpha polypeptide | 3.84 | down | 0.00795 |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| fibrinogen, A alpha polypeptide albumin protease inhibitor 4 (kallistatin) cytochrome P450, subfamily IIE (ethanol-inducible) solute carrier family 2 (facilitated glucose transporter), member 2 EST UDP glycosyltransferase 1 cytochrome P450, subfamily IIIA (niphedipine oxidase), polypeptide 3 carboxypeptidase N, polypeptide 2, 83kD UDP glycosyltransferase 2 family, polypeptide B7 coagulation factor IX (plasma thromboplastic component, Christmas disease, hemophilia B) complement component 9 cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 cytochrome P450, subfamily IVA, polypeptide 11 phypercholesterolemia) cytochrome P450, subfamily IVA, polypeptide 11 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 2, serine hydroxymethyltransferase 1 (soluble) phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- hydroxylase), polypeptide 9 | Affy ID | | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|--|--------------------|---------------|--------|--|-------------|-----------|---------|
| albumin protease inhibitor 4 (kallistatin) 2088 cytochrome P450, subfamily IIE (ethanol-inducible) solute carrier family 2 (facilitated glucose transporter), 2099 member 2 2101 EST 2106 UDP glycosyltransferase 1 cytochrome P450, subfamily IIIA (niphedipine oxidase), polypeptide 3 2117 carboxypeptidase N, polypeptide 2, 83kD 2120 UDP glycosyltransferase 2 family, polypeptide B7 coagulation factor IX (plasma thromboplastic component, 2125 Christmas disease, hemophilia B) 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital-inducible), 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 cytochrome P450, subfamily IIA (aromatic compound- 2131 hypercholesterolemia) cytochrome P450, subfamily IAA, polypeptide 11 2138 cytochrome P450, subfamily IAA, polypeptide 11 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 2, 2150 Coenzyme A ligase, long-chain 2, 2151 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- bydroxylase), polypeptide 9 | HG2730-HT2828_s_at | | | | 3.62 | | 0.01013 |
| HG4533-HT4938 protease inhibitor 4 (kallistatin) J02843 2088 cytochrome P450, subfamily IE (ethanol-inducible) solute carrier family 2 (facilitated glucose transporter), J03810 2099 member 2 J03910 2101 EST J04093 2106 UDP glycosyltransferase 1 cytochrome P450, subfamily IIIA (niphedipine oxidase), J05158 2117 carboxypeptidase N, polypeptide 2, 83kD J05158 2127 carboxypeptidase N, polypeptide 2, 83kD J05168 2127 christmas disease, hemophilia B) K02766 2126 Christmas disease, hemophilia B) K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 cytochrome P450, subfamily I (aromatic compound- L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00352 2132 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00352 2132 polypeptide 6 L04751 2138 cytochrome P450, subfamily I (aromatic compound- L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) L11931 2159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | HG2841-HT2968_s_at | HG2841-HT2968 | | albumin | 4.62 | down | 0.00552 |
| J02843 2088 cytochrome P450, subfamily IIE (ethanol-inducible) solute carrier family 2 (facilitated glucose transporter), J03810 2101 EST J04093 2106 UDP glycosyltransferase 1 cytochrome P450, subfamily IIIA (niphedipine oxidase), J04449 2110 polypeptide 3 J05158 2117 carboxypeptidase N, polypeptide 2, 83kD J05428 2120 UDP glycosyltransferase 2 family, polypeptide BY coagulation factor IX (plasma thromboplastic component, K02402 2125 Christmas disease, hemophilia B) cytochrome P450, subfamily IIA (phenobarbital-inducible), tt K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), tt K03192 2137 polypeptide 6 cytochrome P450, subfamily I (aromatic compound- L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) tt L11931 2159 serine hydroxymethytransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | HG4533-HT4938_at | HG4533-HT4938 | | protease inhibitor 4 (kallistatin) | 3.35 | down | 0.01605 |
| solute carrier family 2 (facilitated glucose transporter), J03810 2099 member 2 J04093 2101 UDP glycosyltransferase 1 Cydchrome P450, subfamily IIIA (niphedipine oxidase), J05428 2120 UDP glycosyltransferase 2 family, polypeptide B7 coagulation factor IX (plasma thromboplastic component, K02402 2125 Christmas disease, hemophilia B) K02766 complement component 9 K03192 2127 polypeptide 6 cydchrome P450, subfamily IIA (phenobarbital-inducible), tr K03192 2127 polypeptide 6 cydchrome P450, subfamily IIA (phenobarbital-inducible), tr L00352 2127 polypeptide 6 cydchrome P450, subfamily I (aromatic compound- 100352 2137 polypeptide 6 cydchrome P450, subfamily I (aromatic compound- 100451 2138 cydchrome P450, subfamily I (aromatic compound- 1004751 2138 cydchrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) tr L1931 2159 serine hydroxymethyltransferase 1 (soluble) cydchrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | J02843_at | J02843 | 2088 | cytochrome P450, subfamily IIE (ethanol-inducible) | 6.18 | down | 0.01308 |
| J03810 2099 member 2 J03910 2101 EST J04493 2106 UDP glycosyltransferase 1 cytochrome P450, subfamily IIIA (niphedipine oxidase), J05428 2120 UDP glycosyltransferase 2 family, polypeptide B7 coagulation factor IX (plasma thromboplastic component, K02402 2125 Christmas disease, hemophilia B) K02766 2126 Christmas disease, hemophilia B) cytochrome P450, subfamily IIA (phenobarbital-inducible), tr K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 cytochrome P450, subfamily I (aromatic compound- 100352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- 1004751 2138 cytochrome P450, subfamily I (aromatic compound- 105144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) tr L1931 2139 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- 116883 2166 hydroxylase), polypeptide 9 | | | | solute carrier family 2 (facilitated glucose transporter), | | | |
| J03910 2101 EST J04093 2106 UDP glycosyltransferase 1 cytochrome P450, subfamily IIIA (niphedipine oxidase), J05428 2110 UDP glycosyltransferase 2 family, polypeptide B7 coagulation factor IX (plasma thromboplastic component, K02402 2125 Christmas disease, hemophilia B) K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), IV K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), IV K03192 2131 hypercholesterolemia) cytochrome P450, subfamily IVA, polypeptide 11 L00352 2131 hypercholesterolemia) cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | J03810_at | J03810 | 2099 | member 2 | 3.6 | down | 0.02376 |
| cytochrome P450, subfamily IIIA (niphedipine oxidase), 10449 2110 polypeptide 3 105428 2117 carboxypeptidase N, polypeptide 2, 83kD 105428 2120 UDP glycosyltransferase 2 family, polypeptide B7 coagulation factor IX (plasma thromboplastic component, CO Christmas disease, hemophilia B) K02766 2125 Christmas disease, hemophilia B) Complement component 9 cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 low density lipoprotein receptor (familial L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- tr L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid- L11931 2159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | | J03910 | 2101 | EST | 18.13 | down | 0.00119 |
| cytochrome P450, subfamily IIIA (niphedipine oxidase), J04449 2110 polypeptide 3 J05158 2117 carboxypeptidase N, polypeptide 2, 83kD J05428 2120 UDP glycosytransferase 2 family, polypeptide B7 coagulation factor IX (plasma thromboplastic component, K02402 2125 Christmas disease, hemophilia B) K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), C10352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid- L09229 2150 Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | J04093_s_at | J04093 | 2106 | UDP glycosyltransferase 1 | 3.28 | down | 0.02286 |
| J04449 2110 polypeptide 3 J05158 2117 carboxypeptidase N, polypeptide 2, 83kD J05428 2120 UDP glycosyltransferase 2 family, polypeptide B7 coagulation factor IX (plasma thromboplastic component, K02402 2125 Christmas disease, hemophilia B) K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 low density lipoprotein receptor (familial L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | | | | cytochrome P450, subfamily IIIA (niphedipine oxidase), | | | |
| J05158 2117 carboxypeptidase N, polypeptide 2, 83kD J05428 2120 UDP glycosyltransferase 2 family, polypeptide B7 coagulation factor IX (plasma thromboplastic component, K02402 2125 Christmas disease, hemophilia B) K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2137 polypeptide 6 low density lipoprotein receptor (familial L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | J04449_at | J04449 | 2110 | polypeptide 3 | 5.07 | down | 0.01733 |
| J05428 2120 UDP glycosyltransferase 2 family, polypeptide B7 coagulation factor IX (plasma thromboplastic component, K02402 2125 Christmas disease, hemophilia B) K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 low density lipoprotein receptor (familial L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L0514 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid- L11931 2159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | J05158_at | J05158 | 2117 | carboxypeptidase N, polypeptide 2, 83kD | 3.37 | down | 0.01156 |
| coagulation factor IX (plasma thromboplastic component, K02402 2125 Christmas disease, hemophilia B) K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital-inducible), cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 low density lipoprotein receptor (familial L00352 2131 hypercholesterolemia) L00389 2132 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound-cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid-coenzyme A ligase, long-chain 2,159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4-chain 2,166 hydroxylase), polypeptide 9 | J05428_at | J05428 | 2120 | UDP glycosyltransferase 2 family, polypeptide B7 | 4.86 | down | 0.03414 |
| K02402 2125 Christmas disease, hemophilia B) K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 low density lipoprotein receptor (familial L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1, fatty-acid- L1931 2159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | | | | coagulation factor IX (plasma thromboplastic component, | | | |
| K02766 2126 complement component 9 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 low density lipoprotein receptor (familial L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 2 L11931 2150 coenzyme A ligase, long-chain 2 L11931 2160 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | K02402_at | K02402 | 2125 | Christmas disease, hemophilia B) | 6.64 | down | 0.04082 |
| cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 low density lipoprotein receptor (familial L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- cytochrome P450, subfamily I (aromatic compound- loud751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid- L11931 2150 Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | K02766_at | K02766 | 2126 | complement component 9 | 5.03 | down | 0.0433 |
| K03192 2127 polypeptide 6 cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 low density lipoprotein receptor (familial L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 2 L11931 2150 Coenzyme A ligase, long-chain 2 L12760 2162 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | | | | cytochrome P450, subfamily IIA (phenobarbital-inducible), | | | |
| Cytochrome P450, subfamily IIA (phenobarbital-inducible), K03192 2127 polypeptide 6 low density lipoprotein receptor (familial L00352 2131 hypercholesterolemia) Cytochrome P450, subfamily I (aromatic compound-cytochrome P450, subfamily IVA, polypeptide 11 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 2 L11931 2150 Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4-L16883 2166 hydroxylase), polypeptide 9 | K03192_f_at | K03192 | 2127 | polypeptide 6 | 14.19 | down | 0.0307 |
| K03192 2127 polypeptide 6 low density lipoprotein receptor (familial L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 2, 150 Coenzyme A ligase, long-chain 2, 159 serine hydroxymethyltransferase 1 (soluble) L12760 2162 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | | | | cytochrome P450, subfamily IIA (phenobarbital-inducible), | | | |
| L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid- L09229 2150 Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | K03192_f_at | K03192 | 2127 | polypeptide 6 | 8.02 | down | 0.03483 |
| L00352 2131 hypercholesterolemia) cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid- L09229 2150 Coenzyme A ligase, long-chain 1,fatty-acid- L11931 2159 serine hydroxymethyltransferase 1 (soluble) L12760 2162 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | | | | low density lipoprotein receptor (familial | | | |
| cytochrome P450, subfamily I (aromatic compound- L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid- L09229 2150 Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) L12760 2162 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | L00352_at | L00352 | 2131 | hypercholesterolemia) | 3.3 | down | 0.03487 |
| L00389 2132 inducible), polypeptide 2 L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid- L09229 2150 Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) L12760 2162 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | | | • | cytochrome P450, subfamily I (aromatic compound- | | | |
| L04751 2138 cytochrome P450, subfamily IVA, polypeptide 11 L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid- L09229 2150 Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) L12760 2162 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | L00389_f_at | F00389 | 2132 | inducible), polypeptide 2 | 3.87 | down | 0.00844 |
| L05144 2139 phosphoenolpyruvate carboxykinase 1 (soluble) fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid- L09229 2150 Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) L12760 2162 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | L04751_at | L04751 | 2138 | cytochrome P450, subfamily IVA, polypeptide 11 | 8.13 | down | 0.02065 |
| fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid-L09229 2150 Coenzyme A ligase, long-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) 2162 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4-L16883 2166 hydroxylase), polypeptide 9 | L05144_at | L05144 | 2139 | phosphoenolpyruvate carboxykinase 1 (soluble) | 4 | down | 0.021 |
| L09229 2150 Coenzyme A ligase, Iong-chain 2 L11931 2159 serine hydroxymethyltransferase 1 (soluble) L12760 2162 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | | | | fatty-acid-Coenzyme A ligase, long-chain 1,fatty-acid- | | | |
| L11931 2159 serine hydroxymethyltransferase 1 (soluble) L12760 2162 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | L09229_s_at | L09229 | 2150 | Coenzyme A ligase, long-chain 2 | 4.5 | down | 0.01347 |
| L12760 2162 phosphoenolpyruvate carboxykinase 1 (soluble) cytochrome P450, subfamily IIC (mephenytoin 4- L16883 2166 hydroxylase), polypeptide 9 | L11931_at | L11931 | 2159 | serine hydroxymethyltransferase 1 (soluble) | 3.74 | down | 0.0056 |
| L16883 2166 hydroxylase), polypeptide 9 | L12760_s_at | L12760 | 2162 | phosphoenolpyruvate carboxykinase 1 (soluble) | 90'9 | down | 0.01005 |
| | L16883_s_at | L16883 | 2166 | hydroxylase), polypeptide 9 | 5.85 | down | 0.04368 |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|-------------|---------|--------|--|-------------|-----------|---------|
| | | | solute carrier family 10 (sodium/bile acid cotransporter | | | |
| L21893 at | L21893 | 2176 | family), member 1 | 5.23 | down | 0.03367 |
| L27050 at | L27050 | 2186 | apolipoprotein F | 4.18 | down | 0.04901 |
| 1.32140_at | L32140 | 2192 | afamin | 5.39 | down | 0.02767 |
| M10942 at | M10942 | 2233 | metallothionein 1E (functional) | 4.05 | down | 0.01412 |
| M10943 at | M10943 | 2234 | metallothionein 1F (functional) | 6.23 | down | 0.00007 |
| M13143_at | M13143 | 2249 | kallikrein B plasma, (Fletcher factor) 1 | 3.04 | down | 0.008 |
| M14777 s at | M14777 | 2263 | glutathione S-transferase A2, glutathione S-transferase A3 | 13.23 | down | 0.03224 |
| M16594 at | M16594 | 2272 | glutathione S-transferase A2 | 5.42 | down | 0.03813 |
| M16750 s at | M16750 | 2273 | pim-1 oncogene | 3.07 | down | 0.02391 |
| M16974 s at | M16974 | 2277 | complement component 8, alpha polypeptide | 10.85 | down | 0.02313 |
| M25079_s_at | M25079 | 2305 | hemoglobin, beta | 4.31 | down | 0.01567 |
| M26393_s_at | M26393 | 2309 | acyl-Coenzyme A dehydrogenase, C-2 to C-3 short chain | 4.3 | down | 0.02294 |
| M29873_s_at | M29873 | 2318 | cytochrome P450, subfamily IIB (phenobarbital-inducible) | 17.92 | down | 0.01469 |
| M29874 s at | M29874 | 2319 | cytochrome P450, subfamily IIB (phenobarbital-inducible) | 8.13 | down | 0.01064 |
| M30185 at | M30185 | 2321 | cholesteryl ester transfer protein, plasma | 3.82 | down | 0.00131 |
| M30185_at | M30185 | 2321 | cholesteryl ester transfer protein, plasma cytochrome P450, subfamily I (aromatic compound- | 3.31 | down | 0.00109 |
| M31667_f_at | M31667 | 2331 | inducible), polypeptide 2 | 4.47 | down | 0.01116 |
| | | | cytochrome P450, subfamily IIA (phenobarbital-inducible), | | | |
| M33317 f at | M33317 | 2338 | polypeptide 7 | 11.47 | down | 0.02611 |
| M34276 at | M34276 | 2341 | plasminogen | 3.08 | down | 0.02754 |
| M57731 s at | M57731 | 2359 | GRO2 oncogene | 3.16 | down | 0.02204 |
| l 1 | | | cytochrome P450, subfamily IIC (mephenytoin 4- | | ٠ | ٠ |
| M61854 s at | M61854 | 2370 | hydroxylase) | 3.45 | down | 0.02949 |
| M63967_at | M63967 | 2378 | aldehyde dehydrogenase 5 | 3.88 | down | 0.00274 |
| M68840 at | M68840 | 2388 | monoamine oxidase A | 3.1 | down | 0.01953 |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank Seq ID | D Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------------|--|-------------|--------------|---------|
| M68895_rna1_at | M68895 2390 |) alcohol dehydrogenase 6 (class V) | 3.21 | down | 0.02095 |
| M72885_rna1_s_at | | | 3.34 | down | 0.02943 |
| M76665_at | | 3 hydroxysteroid (11-beta) dehydrogenase 1 | 90.9 | down | 0.01317 |
| M81349_at | M81349 2405 | 5 serum amyloid A4, constitutive | 10.97 | down | 0.01946 |
| M83652_s_at | M83652 2408 | 3 properdin P factor, complement | 9 | down | 0.00002 |
| M83772_at | M83772 2409 | flavin containing monooxygenase 3 | 5.14 | down | 0.02023 |
| | | insulin-like growth factor binding protein, acid labile | | | |
| M86826_at | | | 3.75 | down | 0.01157 |
| M93405_at | M93405 2424 | t methylmalonate-semialdehyde dehydrogenase | 3.09 | down | 0.03285 |
| M94065_s_at | M94065 2425 | 5 dihydroorotate dehydrogenase | 7.87 | down | 0.0011 |
| M94065_at | M94065 2425 | 5 dihydroorotate dehydrogenase | 3.61 | down | 0.00229 |
| M95585_s_at | M95585 2430 | _ | 3.36 | down | 0.00492 |
| | | transducin-like enhancer of split 4, homolog of Drosophila | | | |
| M99439_at | M99439 2438 | _ | 4.82 | down | 0.00121 |
| rc_N22404_at | N22404 2450 |) EST | 3.44 | down | 0.02267 |
| rc_N22938_s_at | N22938 2453 | serum amyloid A4, constitutive | 4.91 | down | 0.01918 |
| rc N29353 at | N29353 - 2477 | kyniirenine 3-monooxyyaansee (kyniirenine 3-hydroxylasa) | 77 8 | 2000 | 0.04242 |
| rc N29764 at | | | 1 0 | down down | 0.01212 |
| 10 1120101 of | | | † t | nom: | 0.013 |
| rc_N31/41_at | | | 5.66 | down | 0.00212 |
| rc_N34804_at | | | 3.97 | down | 0.0175 |
| rc_N39163_at | | | 4.3 | down | 0.03917 |
| rc_N39201_at | | | 4.79 | down | 0.02015 |
| rc_N49902_at | N49902 2540 | | 3.02 | down | 0.00951 |
| rc_N51117_at | N51117 2544 | | 8.17 | down | 0.00105 |
| rc_N51773_at | N51773 2549 | 9 EST | 6.92 | down | 0.01839 |
| | | LIM protein (similar to rat protein kinase C-binding | | | |
| rc_N52271_at | N52271 2552 | • | 3.67 | down | 0.01102 |
| rc_N52322_at | N52322 2553 | 3 EST | 4.58 | uwop . | 0.02077 |
| rc_N54053_at | N54053 2561 | 1 secreted phosphoprotein 2, 24kD | 12.87 | down | 0.01821 |
| rc_N54417_s_at | N54417 2567 | 7 fibrinogen, A alpha polypeptide | 6.47 | down | 0.00733 |
| | | | | | |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank Seq ID | ID Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|----------------|--|-------------|-----------|---------|
| rc N54429 at | N54429 2 | 2568 EST | 6.85 | down | 0.03334 |
| rc N54950 s at | N54950 28 | 2573 ketohexokinase (fructokinase) | 6.47 | down | 0.0223 |
| N57464 at | N57464 28 | 2576 CCAAT/enhancer binding protein (C/EBP), delta | 4.87 | down | 0.00111 |
| rc N57934 s at | N57934 2577 | _ | 3.28 | down | 0.01555 |
| rc_N58009_at | N58009 28 | 2578 formiminotransferase cyclodeaminase | 8.52 | down | 0.01808 |
| rc N59550 at | N59550 2 | 2588 EST | 4.78 | down | 0.02924 |
| rc N63391 at | N63391 26 | 2600 EST | 3.87 | down | 0.02935 |
| rc N63845 at | N63845 26 | 2605 phytanoyl-CoA hydroxylase (Refsum disease) | 6.82 | down | 0.00369 |
| l I | | enoyl-Coenzyme A, hydratase/3-hydroxyacyl Coenzyme A | ∢ | | |
| rc N64036 s at | N64036 26 | 2607 dehydrogenase | 6.12 | down | 0.00476 |
| rc N65959 at | N65959 2(| 2612 EST | 3.38 | down | 0.00785 |
| rc_N66066_at | N66066 26 | 2613 EST | 4.33 | down | 0.0184 |
| rc_N67105_at | N67105 26 | 2624 EST | 4.69 | down | 0.00194 |
| rc_N68596_s_at | N68596 26 | 2636 betaine-homocysteine methyltransferase | 10.46 | down | 0.01971 |
| rc N70358 s at | N70358 26 | 2657 growth hormone receptor | 8.47 | down | 0.00816 |
| ! ! | | solute carrier family 10 (sodium/bile acid cotransporter | | | |
| rc N70966 s at | N70966 2 | 2663 family), member 1 | 10.8 | down | 0.02894 |
| rc N73543 at | N73543 2 | 2675 EST | 4.64 | down | 0.03981 |
| rc_N74025_at | N74025 20 | 2685 deiodinase, iodothyronine, type l | . 8.18 | down | 0.01363 |
| N77326 at | | 2696 EST | 4.08 | down | 0.00768 |
| rc N80129 i at | | 2703 metallothionein 1L | 26.87 | down | 0.00999 |
| rc_N80129_f_at | N80129 2 | 2703 metallothionein 1L | 11.48 | down | 0.00167 |
| rc N90584 at | N90584 2 | 2714 EST | 3.36 | down | 0.01561 |
| N91087_at | N91087 2 | 2717 EST | 3.66 | down | 0.00725 |
| N99542_at | N99542 2 | 47 orosomucoid 1 | 3.53 | down | 0.00607 |
| rc R01023 s at | R01023 2 | 2752 glucokinase (hexokinase 4) regulatory protein | 4.56 | down | 0.04036 |
| rc_R08564_at | R08564 2 | 2780 plasminogen-like | 8.77 | down | 0.01284 |
| rc_R09053_at | R09053 2 | 2783 EST - | 3.45 | down | 0.03074 |
| rc_R12472_at | R12472 2 | 2789 EST | 12.09 | down | 0.02379 |
| rc_R22905_at | R22905 2 | 2801 EST | 4.31 | down | 0.01744 |
| rc_R40395_s_at | R40395 2 | 2841 lecithin-cholesterol acyltransferase | 12.85 | down | 0.01334 |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank Seq ID | ID Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|----------------|---|-------------|-----------|---------|
| rc_R40492_at | R40492 2842 | t2 EST | 6.4 | down | 0.00527 |
| rc_R40899_f_at | R40899 2844 | - | 4.84 | down | 0.02369 |
| rc_R43799_at | R43799 2851 | 51 EST | 3.93 | down | 0.005 |
| rc_R49602_at | R49602 2885 | | 16.17 | down | 0.00279 |
| rc_R59722_at | R59722 2916 | I6 EST | 6.24 | down | 0.02361 |
| rc_R65593_s_at | | | 9.9 | down | 0.01982 |
| rc_R66002_at | R66002 2936 | _ | 4.33 | down | 0.00789 |
| R69417_at | R69417 2942 | | 6.43 | down | 0.00778 |
| rc_R73816_at | | S1 EST | 7.05 | down | 0.01287 |
| R77628_at | | | 5.51 | down | 0.0404 |
| R79750_at | | | 4.89 | down | 0.00695 |
| R80048_at | R80048 2972 | 72 EST | 3.61 | down | 0.01209 |
| rc_R89811_s_at | | 30 HGF activator | 13.29 | down | 0.00148 |
| rc_R92475_s_at | R92475 2987 | 37 flavin containing monooxygenase 3 | 6.46 | down | 0.02269 |
| rc_R93714_at | R93714 2992 | 32 fetuin B | 4.65 | down | 0.03704 |
| R93776_s_at | R93776 2993 | | 5.55 | down | 0.00084 |
| rc_R94674_s_at | R94674 2996 | 96 EST | 4.58 | down | 0.0047 |
| | | cytochrome P450, subfamily VIIIB (sterol 12-alpha- | | | |
| rc_R97419_at | R97419 3004 | 34 hydroxylase), polypeptide 1 | 19.3 | down | 0.00807 |
| R98073_at | R98073 3009 | 09 EST | 8.37 | down | 0.01436 |
| rc_R99591_at | R99591 3016 | 16 CD5 antigen-like (scavenger receptor cysteine rich family) aldo-keto reductase family 1, member C4 (chlordecone reductase: 3-aloha hydroxysteroid dehydrogenase, tyne I: | 7.41 | down | 0.00043 |
| S68287 at | S68287 3025 | _ | 5.04 | down | 0.02895 |
| S70004_at | | - | 5.13 | down | 0.00183 |
| S77356_at | | | 3.55 | down | 0.03874 |
| rc_t10264_s_at | | | 3.26 | down | 0.01718 |
| rc_T16484_s_at | | | 4.78 | down | 60000.0 |
| rc_T40936_at | T40936 3118 | 18 EST | 4.62 | down | 0.02844 |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| rc_T87174_at | T87174 | 3240 | EST | 3.46 | down | 0.00026 |
| T95813_f_at | T95813 | 3262 | KIAA1051 protein | 20.36 | down | 0.01361 |
| rc_T98199_i_at | T98199 | 3267 | EST | 4.05 | down | 0.00753 |
| rc_T98676_at | T98676 | 3269 | EST | 11.15 | down | 0.0323 |
| U02388_at | U02388 | 3278 | cytochrome P450, subfamily IVF, polypeptide 2 | 4.4 | down | 0.00761 |
| U06641_s_at | U06641 | 3287 | UDP glycosyltransferase 2 family, polypeptide B15 | 6.37 | down | 0.01594 |
| U08006_s_at | 00800 | 3290 | complement component 8, alpha polypeptide | 3.96 | down | 0.04272 |
| U08021_at | U08021 | 3291 | nicotinamide N-methyltransferase | 3.63 | down | 0.03726 |
| U20530_at | U20530 | 3322 | secreted phosphoprotein 2, 24kD | 5.31 | down | 0.01119 |
| U21931_at | U21931 | 3326 | fructose-bisphosphatase 1 | 3.17 | down | 0.0143 |
| | | | cytochrome P450, subfamily IIA (phenobarbital-inducible), | | | |
| U22029_f_at | U22029 | 3327 | polypeptide 7 | 11.85 | down | 0.03538 |
| | | | solute carrier family 6 (neurotransmitter transporter, | | | |
| U27699_at | U27699 | 3340 | betaine/GABA), member 12 | 3.65 | down | 0.00381 |
| U50196_at | U50196 | 3377 | adenosine kinase | 3.03 | down | 0.00975 |
| U50929_at | U50929 | 3380 | betaine-homocysteine methyltransferase | 8.04 | down | 0.0188 |
| U51010_s_at | U51010 | 3381 | nicotinamide N-methyltransferase | 4.69 | down | 0.03099 |
| U56814_at | U56814 | 3393 | deoxyribonuciease I-like 3 | 17.69 | down | 0.00007 |
| U56814_at | U56814 | 3393 | deoxyribonuclease I-like 3 | 5.75 | down | 0.00152 |
| U65932_at | U65932 | 3405 | extracellular matrix protein 1 | 3.18 | down | 0.00575 |
| U95090_at | N95090 | 3464 | nephrosis 1, congenital, Finnish type (nephrin) | 4.63 | down | 0.01595 |
| W07723_at | W07723 | 3471 | EST | 3.51 | down | 0.00026 |
| W26996_at | W26996 | 3484 | EST | 4.46 | down | 0.00734 |
| W28414_at | W28414 | 3490 | EST | 4.06 | down | 0.00083 |
| W28798_at | W28798 | 3492 | phosphodiesterase 6A, cGMP-specific, rod, alpha | 3.33 | down | 0.00222 |
| W28944_at | W28944 | 3494 | EST | 6.9 | down | 0.01014 |
| rc_W44745_at | W44745 | 3520 | EST | 3.87 | down | 0.01051 |
| rc_W45560_at | W45560 | 3525 | EST | 3.48 | down | 0.0179 |
| W55903_at | W55903 | 3546 | adipose differentiation-related protein; adipophilin | 5.64 | down | 0.00014 |
| rc_W63728_at | W63728 | 3565 | EST | 3.86 | down | 0.00288 |
| rc_W67147_at | W67147 | 3568 | deleted in liver cancer 1 | 4.37 | down | 0.00069 |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| rc_W72044_at rc_W72382_at rc_W73601_at | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|--|---------|--------|---|-------------|-----------|---------|
| rc_W72382_at rc_W73601_at | W72044 | 3580 | insulin induced gene 1 | 3.1 | down | 0.03445 |
| rc_W72382_at rc_W73601_at | | | oxidative 3 alpha hydroxysteroid denydrogenase; retinol | | | |
| rc_W73601_at | W72382 | 3584 | dehydrogenase | 68.6 | down | 0.03091 |
| 1 | W73601 | 3592 | EST | 3.45 | down | 0.01382 |
| rc W73818 at | W73818 | 3593 | EST | 3.47 | down | 0.00927 |
| rc_W81552_at | W81552 | 3615 | EST | 12.97 | down | 0.00244 |
| rc_W86075_at | W86075 | 3624 | EST | 6.04 | down | 0.01486 |
| rc_W86600_at | W86600 | 3628 | EST | 3.67 | down | 0.04208 |
| rc_W87532_at | W87532 | 3634 | putative glycine-N-acyltransferase | 5.5 | down | 0.00739 |
| rc_W87781_at | W87781 | 3636 | EST | 4.02 | down | 0.00284 |
| rc_W88946_at | W88946 | 3639 | putative glycine-N-acyltransferase | 25.28 | down | 0.00221 |
| rc_W95041_at | W95041 | 3662 | EST | 4.22 | down | 0.01005 |
| X02176 s at | X02176 | 3672 | complement component 9 | 3.84 | down | 0.01793 |
| | X06562 | 3686 | growth hormone receptor | 4.8 | down | 0.00507 |
| X06985_at | X06985 | 3689 | heme oxygenase (decycling) 1 | 3.34 | down | 0.00045 |
| X13227 at | X13227 | 3698 | D-amino-acid oxidase | 3.22 | down | 0.01753 |
| I | | | cytochrome P450, subfamily IIA (phenobarbital-inducible), | | | |
| X13930 f at | X13930 | 3700 | polypeptide 6 | 8.1 | down | 0.0219 |
| i I | | | acetyl-Coenzyme A acyltransferase 1 (peroxisomal 3- | | | |
| X14813 at | X14813 | 3705 | oxoacyl-Coenzyme A thiolase) | 3.53 | down | 0.00059 |
| X16260 s at | X16260 | 3710 | inter-alpha (globulin) inhibitor, H1 polypeptide | 3.76 | down | 0.00291 |
| X16349 s at | X16349 | 3712 | sex hormone-binding globulin | 6.61 | down | 0.00008 |
| X54380 at | X54380 | 3730 | pregnancy-zone protein | 7.71 | down | 0.00069 |
| X56411 rna1 at | X56411 | 3737 | alcohol dehydrogenase 4 (class II), pi polypeptide | 9.87 | down | 0.01416 |
| X58022 at | X58022 | 3747 | corticotropin releasing hormone-binding protein | 4.09 | Howop | 0.00076 |
| X63359_at | X63359 | 3759 | UDP glycosyltransferase 2 family, polypeptide B10 | 4.26 | down | 0.01725 |
| X64177 f at | X64177 | 3763 | metallothionein 1H | 3.26 | down | 0.03928 |
| X67491 f at | X67491 | 3776 | glutamate dehydrogenase 1 | 4.06 | down | 0.00273 |
| X72177 rna1 at | X72177 | 3790 | complement component 6 | 4.25 | down | 0.01598 |
| X76717 at | X76717 | 3797 | metallothionein 1L | 5.64 | down | 0.00215 |
| X90579 s at | X90579 | 3819 | EST | 4.26 | down | 0.04759 |

Table 7A. Up reglated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change Direction Pvalue | Direction | Pvalue |
|------------------|---------|--------|--|------------------------------|-----------|---------|
| X95190_at | X95190 | 3832 | acyl-Coenzyme A oxidase 2, branched chain | 6.22 | down | 0.00162 |
| X97324_at | X97324 | 3839 | adipose differentiation-related protein; adipophilin | 3.72 | down | 0.00202 |
| Y00317_at | Y00317 | 3845 | UDP glycosyltransferase 2 family, polypeptide B4 | 4.63 | down | 0.02986 |
| Z20777_at | Z20777 | 3866 | EST | 15.73 | down | 0.00147 |
| l | | | aldo-keto reductase family 1, member D1 (delta 4-3- | | | |
| Z28339_at | Z28339 | 3875 | ketosteroid-5-beta-reductase) | 8.03 | down | 0.00853 |
| | | | lipase A, Iysosomal acid, cholesterol esterase (Wolman | | | |
| Z31690_s_at | Z31690 | 3881 | disease) | 3.29 | down | 0.00161 |
| rc Z40259 s at | Z40259 | 3919 | EST | 4.47 | down | 0.00093 |
| rc_Z40305_at | Z40305 | 3920 | EST | 4.09 | down | 96000.0 |
| rc_Z40902_at | Z40902 | 3926 | SEC14 (S. cerevisiae)-like 2 | 4.97 | down | 0.04627 |
| rc_Z41042_at | Z41042 | 3928 | EST | 3.37 | down | 0.00703 |
| Z48475 at | Z48475 | 3943 | glucokinase (hexokinase 4) regulatory protein | 4.6 | down | 0.01693 |
| | | | small inducible cytokine subfamily A (Cys-Cys), member | | | |
| Z49269_at | Z49269 | 3945 | 14 | 7.24 | down | 0.01047 |
| Z69923_at | Z69923 | 3948 | HGF activator | 3.95 | down | 0.00012 |
| | | | | | | |
| Z80345_rna1_s_at | Z80345 | 3951 | acyl-Coenzyme A dehydrogenase, C-2 to C-3 short chain | 3.21 | down | 0.04734 |
| Z84721_cds2_at | Z84721 | 3953 | hemoglobin, zeta | 7.39 | down | 0.01921 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| rc AA001504 f at | AA001504 | 2 | EST | 4.44 | dn | 0.03077 |
| rc_AA005262_at | AA005262 | 13 | EST | 3.09 | 롸 | 0.0064 |
|] | | | KIAA1080 protein; Golgi-associated, gamma-adaptin ear | | | |
| rc AA007507 at | AA007507 | 18 | containing, ARF-binding protein 2 | 5.23 | dn | 0.00159 |
| rc AA010065 s at | AA010065 | 22 | CDC28 protein kinase 2 | 3.71 | d | 0.00432 |
| rc_AA011209_s_at | AA011209 | 30 | melanoma-associated antigen recognised by T | 6.45 | dn | 0.00088 |
| rc AA011679 at | AA011679 | 32 | EST | 3.08 | dn | 0.03649 |
| rc_AA018346_at | AA018346 | 38 | EST | 3.69 | dn | 0.04582 |
| rc_AA021549_at | AA021549 | 42 | EST | 3.17 | ф | 0.00158 |
| rc_AA022623_at | AA022623 | 4 | EST | 3.27 | dn | 0.01556 |
| rc_AA024658_at | AA024658 | 47 | ribosomal protein S19 | 7.55 | dn | 0.00592 |
| rc_AA024776_at | AA024776 | 48 | EST | 3.44 | dn | 0.00334 |
| rc AA025166 s at | AA025166 | 20 | fusion, derived from t(12;16) malignant liposarcoma | 3.17 | dn | 600000 |
| rc AA026356 at | AA026356 | 24 | EST | 5.04 | dn | 0.02483 |
| rc AA027833 i at | AA027833 | 26 | EST | 5.02 | ф | 0.01123 |
| rc_AA029288_at | AA029288 | 92 | EST | 3.36 | dn | 0.04908 |
| rc_AA031814_at | AA031814 | 2 | KIAA0958 protein | 3.07 | d | 0.00681 |
| rc AA037058 s at | AA037058 | 84 | laminin, gamma 1 (formerly LAMB2) | 4.11 | dn | 0.02264 |
| rc AA037433 at | AA037433 | 86 | EST | 4.9 | dn | 0.0194 |
| rc_AA037766_at | AA037766 | 87 | EST | 3.63 | ф | 0.0328 |
| rc_AA040465_at | AA040465 | 95 | EST | 3.63 | dn | 0.01806 |
| AA043111 s at | AA043111 | 97 | EST | 6.36 | g | 0.0005 |
| rc AA043959_at | AA043959 | 101 | tropomyosin 4 | 4.37 | ф | 0.01641 |
| rc_AA045365_at | AA045365 | 106 | EST | 3.17 | dn | 0.0149 |
| rc_AA046103_at | AA046103 | 109 | EST | 3.75 | dn | 0.02893 |
| rc AA046410 s at | AA046410 | 110 | EST | 3.18 | dn | 0.00797 |
| rc AA046745 at | AA046745 | 113 | Wolf-Hirschhorn syndrome candidate 1 | 3.33 | dn | 0.00648 |
| rc AA047379 s at | AA047379 | 119 | karyopherin (importin) beta 1 | 3.15 | g | 0.01572 |
| rc AA047704 at | AA047704 | 120 | EST | 3.2 | dn | 0.0029 |
| rc_AA052941_at | AA052941 | 121 | EST | 3.36 | dn | 0.00088 |
| rc_AA053662_f_at | AA053662 | 129 | EST | 3.3 | ౚ | 0.00558 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| 00 at 6. at 7. at | 4A053680 | 130 | | 4.07 | ٤ | 0 03144 |
|---|-----------------|-----|---|-------|--------|---------|
| , , , , , , , , , , , , , , , , , , , | AAREBOO | | high-mobility group protein 2-like 1 | 5 | 3 7 | 2 |
| , # , # , # , # , # , # | 760000 | 134 | EST | 3.02 | d | 0.04984 |
| _#_#_# _#_#_# | AA055896 | 135 | collagen, type V, alpha 1 | 10.87 | d | 0.00907 |
| at at | AA070206 | 155 | EST | 3.15 | d | 0.03914 |
| jat Jat | AA070485 | 156 | interleukin 13 receptor, alpha 1 | 3.19 | d | 0.03465 |
| | AA070827 | 157 | EST | 4.37 | dn | 0.02617 |
| AA071387_at A | AA071387 | 158 | jumping translocation breakpoint | 3.31 | dn | 0.0001 |
| , # | AA074162 | 159 | superkiller viralicidic activity 2 (S. cerevisiae homolog)- | 3.23 | dn | 0.00642 |
| rc_AA076138_at A | AA076138 | 167 | H2A histone family, member Y | 3.75 | dn | 0.01442 |
| • | AA086071 | 184 | chromosome-associated polypeptide C | 3.77 | dn | 0.01993 |
| | | | kangai 1 (suppression of tumorigenicity 6, prostate; | | | |
| | | | CD82 antigen (R2 leukocyte antigen, antigen detected | | | |
| fat | AA086232 | 186 | by monoclonal and antibody IA4)) | 4.52 | dn | 0.00452 |
| rc_AA086412_at A | AA086412 | 187 | DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 16 | 3.13 | dn | 0.00327 |
| AA089997_at A | AA089997 | 189 | EST | 4.9 | dn | 0.0241 |
| at | 4A091752 | 193 | purine-rich element binding protein B | 3.25 | dn | 0.01419 |
| , at | 4A092129 | 194 | EST | 2.67 | dn | 0.00011 |
| at | AA092290 | 195 | EST | 3.25 | dn | 0.01616 |
| at | AA094752 | 203 | hypothetical 43.2 Kd protein | 3.44 | dn | 0.04445 |
| ` # | AA099404 | 208 | EST | 20.22 | dn | 0 |
| , ;; | AA101272 | 215 | EST | 3.83 | dn | 0.0386 |
| at | AA102489 | 219 | EST | 5.28 | dn | 0.02122 |
| f_at | AA102837 | 221 | EST | 4.13 | dn | 0.0067 |
| at | AA112679 | 224 | EST | 4.19 | dn | 0.00572 |
| at at | AA115562 | 229 | EST | 3.35 | . dn | 0.00283 |
| ` # | AA115735 | 230 | EST | 4.8 | dn | 0.02671 |
| at | AA116036 | 233 | chromosome 20 open reading frame 1 | 3.41 | dn | 0.00089 |
| | AA122386 | 239 | collagen, type V, alpha 2 | 3.44 | dn | 0.02566 |
| • | 4A125808 | 240 | EST | 3.04 | đ | 0.02112 |
| | 2A127444 | 252 | EST | 3.87 | dn | 0.01751 |
| rc_AA127741_at A | AA127741 | 256 | EST | 4.49 | g n | 0.0463 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| rc AA128407 at | AA128407 | 259 | EST | 3.33 | dn | 0.02298 |
| rc_AA129757_at | AA129757 | 264 | EST | 3.75 | ф | 0.0166 |
| rc_AA131220_at | AA131220 | 267 | EST | 3.18 | dn | 0.00974 |
| rc AA132032 s at | AA132032 | 271 | trinucleotide repeat containing 1 | 3.84 | ф | 0.01136 |
| rc AA132514 at | AA132514 | 272 | EST | 3.2 | ф | 0.00876 |
| rc_AA133527_at | AA133527 | 281 | EST | 5.23 | ф | 0.00037 |
| rc_AA133666_s_at | AA133666 | 283 | cysteine-rich protein 2 | 5.35 | ф | 0.00433 |
| rc_AA134052_s_at | AA134052 | 285 | Rab geranylgeranyltransferase, alpha subunit | 5.47 | dn | 0.00982 |
| rc_AA135153 at | AA135153 | 291 | EST | 5.58 | dn | 0.00327 |
| rc_AA135871_at | AA135871 | 294 | EST | 3.56 | dn | 0.01718 |
| rc_AA136269_at | AA136269 | 298 | EST | 7.5 | ф | 0.00014 |
| rc_AA136474_at | AA136474 | 301 | Meis (mouse) homolog 2 | 3.15 | dn | 0.02837 |
| rc_AA136547_at | AA136547 | 305 | EST | 4.19 | d | 0.00098 |
| rc_AA136864_at | AA136864 | 304 | zinc finger protein homologous to Zfp-36 in mouse | 3.31 | dn | 0.00346 |
| rc_AA142857_at | AA142857 | 307 | EST | 9.48 | 롸 | 0.00376 |
| rc_AA142858_at | AA142858 | 308 | EST | 4.07 | d | 0.0022 |
| rc AA146849 s at | AA146849 | 313 | tärget of myb1 (chicken) homolog | 4.72 | 롸 | 0.00326 |
| rc AA148885 at | AA148885 | 320 | minichromosome maintenance deficient (S. cerevisiae) 4 | 6.59 | 롸 | 0.00112 |
| rc_AA148977_at | AA148977 | 322 | EST | 9.3 | 롸 | 0.00002 |
| rc_AA149889_at | AA149889 | 326 | neighbor of A-kinase anchoring protein 95 | 8.55 | dn | 0.00224 |
| rc_AA151435_at | AA151435 | 336 | EST | 4.52 | 롸 | 0.01134 |
| | | | ATP synthase, H+ transporting, mitochondrial F0 | | | |
| rc_AA156187_at | AA156187 | 339 | complex, subunit b, isoform 1 | 9.38 | 롸 | 0.02007 |
| rc_AA156460_at | AA156460 | 343 | EST | 4.39 | 롸 | 0.01223 |
| rc_AA159025 at | AA159025 | 353 | EST | 6.58 | 롸 | 0.01946 |
| rc_AA160775_s_at | AA160775 | 322 | BCL2-antagonist of cell death | 3.8 | 음 | 0.01145 |
| | | | 3-prime-phosphoadenosine 5-prime-phosphosulfate | | | |
| rc_AA165526_at | AA165526 | 360 | synthase 1 | 3.68 | 음 | 0.00021 |
| rc_AA167708_at | AA167708 | 363 | EST | 3.19 | 롸 | 0.01871 |
| rc_AA171760_at | AA171760 | 367 | EST | 4.39 | 롸 | 0.04582 |
| rc_AA173430_at | AA173430 | 371 | EST | 3.74 | 롸 | 0.01159 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| AA173505_at | AA173505 | 372 | EST | က | G | 0.01736 |
| AA173597_at | AA173597 | 373 | EST | 3.37 | d | 0.03622 |
| rc_AA173755_at | AA173755 | 374 | EST | 6.73 | d | 0.00666 |
| rc_AA179787_at | AA179787 | 380 | polyglutamine binding protein 1 | 4.71 | 9 | 0.00725 |
| rc_AA179845_at | AA179845 | 381 | EST | 3.55 | g | 0.02484 |
| rc_AA181580_s_at | AA181580 | 383 | karyopherin (importin) beta 1 | 3.01 | g. | 0.0125 |
| rc_AA181705_f_at | AA181705 | 385 | EST | 5.9 | g | 0.00023 |
| rc_AA182001_r_at | AA182001 | 386 | EST | 3.78 | 9 | 0.04446 |
| AA187579_at | AA187579 | 330 | MCT-1 protein | 3.4 | g. | 0.02455 |
| rc_AA188378_i_at | AA188378 | 392 | EST | 4.88 | 음 | 0.01653 |
| rc_AA194730_at | AA194730 | 410 | EST | 4.57 | dn | 0.00801 |
| rc_AA194998_at | AA194998 | 413 | purinergic receptor (family A group 5) | 3.06 | a | 0.04752 |
| rc_AA195067_i_at | AA195067 | 414 | GTPase activating protein-like | 3.24 | dn | 0.00606 |
| rc_AA204927_at | AA204927 | 425 | tropomyosin 1 (alpha) | 6.11 | d | 0.0014 |
| rc_AA207103_at | AA207103 | 429 | EST | 3.36 | g | 0.00131 |
| rc_AA211483_at | AA211483 | 435 | EST | 4.11 | d | 0.0365 |
| AA215299_s_at | AA215299 | 439 | U6 snRNA-associated Sm-like protein LSm7 | 4.81 | dn | 0.00119 |
| rc_AA215379_at | AA215379 | 440 | EST | 4.44 | g | 0.01675 |
| rc_AA218663_at | AA218663 | 444 | acid-inducible phosphoprotein | 4.34 | 음 | 0.00161 |
| rc_AA226932_at | AA226932 | 453 | DKFZP564F0923 protein | 5.25 | 9- | 0.00612 |
| rc_AA227145_at | AA227145 | 454 | EST | 3.4 | 읔 | 0.03422 |
| rc_AA227541_at | AA227541 | 457 | NS1-binding protein | 3.6 | 合 | 0.02801 |
| AA232837_at | AA232837 | 465 | EST | 8.85 | 음 | 0.0048 |
| rc_AA233897_at | AA233897 | 476 | EST | 3.8 | 음 | 0.02145 |
| rc_AA234096_at | AA234096 | 479 | EST | 5.75 | S | 0.01169 |
| rc_AA235289_at | AA235289 | 495 | RAP2A, member of RAS oncogene family | 4.31 | g | 0.00135 |
| AA235448_s_at | AA235448 | 497 | EST | 5.62 | d | 0.00077 |
| rc_AA235853_at | AA235853 | 503 | CGI-96 protein | 3.16 | 읔 | 0.00744 |
| rc_AA235868_at | AA235868 | 504 | nuclear transcription factor Y, beta | 3.49 | g | 0.01897 |
| | | | 3-prime-phosphoadenosine 5-prime-phosphosulfate | | | |
| _AA236150_at | AA236150 | 207 | synthase 1 | 3.46 | 음 | 0.0008 |
| rc_AA236150_at | AA236150 | 202 | synthase 1 | | 3.46 | |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| AA236412 at | AA236412 | 511 | EST | 3.1 | dn | 0.04463 |
| rc AA236532 s at | AA236532 | 513 | EST | 3.04 | ф | 0.03747 |
| rc AA236672 at | AA236672 | 515 | EST | 4.37 | 合 | 0.00385 |
| rc_AA236904_at | AA236904 | 518 | EST | 3.07 | dn | 0.01503 |
| | AA242757 | 522 | EST | 3.27 | dn | 0.00286 |
| rc AA243133 at | AA243133 | 525 | serine/threonine kinase 15 | 7.03 | dn | 0.00005 |
| rc_AA243173_at | AA243173 | 526 | EST | 3.49 | dn | 0.0401 |
| AA249819 s at | AA249819 | 535 | EST | 5.22 | dn | 0.00049 |
| rc AA251230 at | AA251230 | 540 | EST | 3.25 | d | 0.01417 |
| rc AA251299 s at | AA251299 | 541 | KIAA0014 gene product | 4.74 | dn | 0.0252 |
| rc AA251428 at | AA251428 | 542 | DKFZP586I2223 protein | 3.15 | ф | 0.01223 |
| rc_AA251766_at | AA251766 | 543 | EST | 3.06 | dn | 0.0098 |
| rc_AA251769_at | AA251769 | 544 | EST | 4.45 | d | 0.01431 |
| rc_AA251792_at | AA251792 | 546 | fatty-acid-Coenzyme A ligase, long-chain 4 | 7.44 | ф | 0.00285 |
| rc_AA251909_at | AA251909 | 549 | EST | 3.59 | dn | 0.01129 |
| rc_AA252060_at | AA252060 | 550 | EST | 4.88 | ф | 0.00169 |
| rc_AA252355_at | AA252355 | 553 | EST | 3.02 | d | 0.00715 |
| rc_AA252524_at | AA252524 | 555 | EST | 3.17 | dn | 0.00686 |
| 1 I | | | chaperonin containing TCP1, subunit 6A (zeta 1), homeo | | | |
| rc AA252627 s at | AA252627 | 556 | box B5 | 4.28 | dn | 0.00363 |
| | AA253011 | 558 | KIAA0713 protein | 3.15 | dn | 0.00035 |
| rc_AA255486 at | AA255486 | 568 | EST | 3.72 | d | 0.00154 |
| rcAA256131_at | AA256131 | 574 | glycophosphatidylinositol anchor attachment 1 | 3.16 | dn | 0.00011 |
| rc_AA256268_at | AA256268 | 216 | EST | 3.13 | dn | 0.03874 |
| rc_AA256524_at | AA256524 | 580 | AD022 protein | 3.06 | dn | 0.00626 |
| rc_AA256606_at | AA256606 | 581 | EST | 3.92 | dn | 0.03087 |
| rc AA256688 s at | AA256688 | 584 | EST | 4.23 | dn | 0.03094 |
| rc AA258131 at | AA258131 | 287 | putative GTP-binding protein similar to RAY/RAB1C | 6.23 | dn | 0.00931 |
| rc_AA258182_at | AA258182 | 589 | EST | 3.55 | d | 0.01198 |
| rc_AA258387_at | AA258387 | 594 | EST | 3.15 | d | 0.02028 |
| rc_AA258421_at | AA258421 | 595 | hypothetical protein | 6.5 | đ | 0.00559 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank Se | Seq ID Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|--------------|---|-------------|-----------|---------|
| rc_AA258614_s_at | AA258614 5 | 599 EST | 3.94 | 유 | 0.0048 |
| rc_AA262477_at | AA262477 6 | 608 ribonuclease HI, large subunit | 4.57 | 롸 | 0.00724 |
| rc_AA262957_at | AA262957 6 | 612 EST | 3.76 | 음 | 0.00157 |
| | | ATP synthase, H+ transporting, mitochondrial F0 | F0 | | |
| AA263032_s_at | AA263032 614 | 4 complex, subunit b, isoform 1 | 6.73 | 9 | 0.04478 |
| rc_AA278768_f_at | AA278768 617 | | 3.77 | 음 | 0.03239 |
| rc_AA278817_at | , | _ | 3.5 | đ | 0.01159 |
| rc_AA279418_at | AA279418 6 | 626 EST | 3.23 | g | 0.02054 |
| rc_AA280734_i_at | AA280734 639 | 9 KIAA0618 gene product | 6.83 | 음 | 0.001 |
| rc_AA280840_at | AA280840 641 | _ | 4.51 | đ | 0.0186 |
| rc_AA281599_at | AA281599 647 | | 4.87 | dn | 0.00248 |
| rc_AA282247_at | AA282247 657 | 7 EST - | 5.88 | dn | 0.01112 |
| rc_AA282343_at | | _ | 5.78 | dn | 0.00128 |
| rc_AA282571_at | | _ | 3.16 | dn | 0.01355 |
| rc_AA283182_at | AA283182 668 | _ | 6.78 | dn | 0.01784 |
| rc_AA283832_at | AA283832 6 | | 4.77 | d | 0.00156 |
| rc_AA284565_f_at | | | 3.27 | ф | 0.0362 |
| rc_AA284720_at | | _ | 3.03 | dn | 0.00252 |
| rc_AA284945_at | AA284945 6 | 680 EST . | 6.25 | 함 | 0.0002 |
| rc_AA285132_at | AA285132 682 | •• | 3.1 | 롸 | 0.00844 |
| rc_AA286911_at | | 684 EST | 3.36 | đ | 0.00037 |
| rc_AA291137_at | AA291137 694 | | 3.67 | dn | 0.03243 |
| rc_AA291139_at | AA291139 6 | _ | 6.22 | 함 | 0.03491 |
| rc_AA291168_at | AA291168 6 | | 4.93 | ᅀ | 0.01633 |
| rc_AA291644_at | | | 3.28 | 함 | 0.00033 |
| rc_AA291659_at | | <u> </u> | 4.15 | dn | 0.00019 |
| AA291786_s_at | • | _ | 4.15 | dn | 0.00362 |
| rc_AA292765_at | AA292765 7 | | 7.24 | dn | 0.00498 |
| rc_AA292788_s_at | AA292788 7 | _ | 3.65 | ďn | 0.01765 |
| rc_AA293420_s_at | | 717 EST | 4.05 | ф | 0.01189 |
| rc_AA293589_s_at | AA293589 7 | 719 zinc finger protein | 3.02 | ф | 0.01809 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| AA293868 s at | AA293868 | 721 | EST | 3.04 | 함 | 0.0054 |
| ຸທ | AA296994 | 724 | seven transmembrane domain protein | 3.16 | ф | 0.0076 |
| AA313213 at | AA313213 | 732 | flotillin 1 | 3.59 | dn | 0.00878 |
| AA320369 s at | AA320369 | 735 | chromosome 19 open reading frame 3 | 3.88 | g | 0.00452 |
| rc AA321833 at | AA321833 | 736 | EST | 3.16 | dn | 0.00523 |
| | AA335191 | 741 | creatine kinase, brain | 6.47 | dn | 0.01462 |
| rc AA338760 at | AA338760 | 744 | EST | 3.96 | dn | 0.01307 |
| rc AA365708 s at | AA365708 | 764 | microfibrillar-associated protein 1 | 3.01 | dn | 0.02372 |
| AA365742 s at | AA365742 | 765 | tetraspan NET-6 protein | 4.12 | ďn | 0.00255 |
| rc AA370163 at | AA370163 | 992 | EST | 3.41 | ф | 0.00134 |
| AA384184 s at | AA384184 | 774 | DKFZP586B0519 protein | 3.42 | dn | 0.01222 |
| AA393139 at | AA393139 | 775 | geminin | 7.44 | ф | 0.00888 |
| rc AA394258 s at | AA394258 | 779 | RD RNA-binding protein | 7.27 | dn | 0.00054 |
| rc AA398141 at | AA398141 | 788 | EST | 3.3 | dn | 0.00211 |
| rc_AA398205_at | AA398205 | 789 | EST | 4.22 | dn | 0.00059 |
| rc_AA398563_at | AA398563 | 797 | EST | 3,14 | dn | 0.01895 |
| rc_AA398908_at | AA398908 | 801 | EST . | 20.72 | dn | 0.00114 |
| rc AA398926 f at | AA398926 | 802 | EST | 8.25 | ф | 0.00066 |
| rc_AA399251_at | AA399251 | 804 | EST | 4.3 | 함 | 0.01578 |
| rc_AA399264_at | AA399264 | 805 | EST | 3.51 | dn | 0.00327 |
| rc_AA400184_at | AA400184 | 808 | KIAA0907 protein | 4.11 | dn | 0.01123 |
| AA400643 s at | AA400643 | 817 | GAS2-related on chromosome 22 | 4.04 | dn | 0.03751 |
| rc AA400896 at | AA400896 | 822 | EST | 3.54 | dn | 0.00889 |
| rc_AA401965_at | AA401965 | 833 | tumor suppressor deleted in oral cancer-related 1 | 7.58 | dn | 0.00089 |
| rc_AA402272_at | AA402272 | 837 | EST | 3.73 | d | 0.02336 |
| rc_AA402968_at | AA402968 | 844 | EST | 3.68 | dn | 0.00123 |
| 1 | | | O-linked N-acetylglucosamine (GlcNAc) transferase | | | |
| | | | (UDP-N-acetylglucosamine:polypeptide-N- | • | | ! |
| rc_AA404560_at | AA404560 | 853 | acetylglucosaminyl transferase) | 3.73 | dn | 0.0143 |
| rc_AA405098_at | AA405098 | 855 | EST | 6.09 | g | 0.01224 |
| rc_AA405505_at | AA405505 | 860 | RNA helicase family | 4.05 | 음 | 0.00/4/ |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|-------------------|----------|--------|---|-------------|-----------|---------|
| rc_AA405544_f_at | AA405544 | 861 | EST | 3.09 | ds S | 0.04146 |
| rc_AA405791_at | AA405791 | 864 | EST | 11.79 | d | 0.00587 |
| rc_AA406216_at | AA406216 | 871 | EST | 3.4 | g. | 0.00529 |
| rc_AA406384_at | AA406384 | 875 | KIAA0670 protein/acinus | 3.23 | 9 | 0.00486 |
| rc_AA410469_at | AA410469 | 883 | EST | 5.45 | d n | 0.00068 |
| rc_AA410962_s_at | AA410962 | 887 | peroxisome proliferative activated receptor, delta | 4.91 | dn | 0.0044 |
| rc_AA412301_at | AA412301 | 899 | EST | 3.42 | ф | 0.0129 |
| rc_AA412720_at | AA412720 | 902 | EST | 3.06 | ф | 0.02153 |
| rc_AA416970_at | AA416970 | 912 | Mad4 homolog | 5.3 | dn | 0.00418 |
| rc_AA416973_at | AA416973 | 913 | EST | 4.29 | dn | 0.00155 |
| rc_AA417030_at | AA417030 | 914 | EST | 7.35 | d | 0.00555 |
| rc_AA417884_at | AA417884 | 919 | cyclin-dependent kinase inhibitor 2C (p18, inhibits | 3.42 | 슘 | 0.02997 |
| AA421213_at | AA421213 | 931 | Lsm3 protein | 3.34 | dn | 0.00198 |
| rc_AA421562_at | AA421562 | 934 | anterior gradient 2 (Xenepus laevis) homolog | 5.02 | 육 | 0.02818 |
| rc_AA421951_at | AA421951 | 936 | EST | 69.9 | ф | 0.00013 |
| rc_AA423827_f_at | AA423827 | 941 | chromosome 22 open reading frame 3 | 4.39 | ф | 0.00345 |
| rc_AA423841_f_at | AA423841 | 942 | EST | 3.71 | d | 0.01481 |
| rc_AA424029_at | AA424029 | 943 | EST | 4.54 | dn | 0.02721 |
| rc_AA424487_at | AA424487 | 945 | EST | 4.68 | g n | 0.0013 |
| rc_AA424881_at | AA424881 | 949 | EST | 3.39 | g | 0.03546 |
| | | | eukaryotic translation initiation factor 2B, subunit 2 (beta, | | | |
| rc_AA425544_s_at | AA425544 | 955 | 39kD) | 3.05 | 음 | 0.0346 |
| rc_AA425852_s_at, | AA425852 | 928 | EST | 3.98 | dn | 0.02796 |
| rc_AA425852_l_at | AA425852 | 928 | EST | 3.82 | g | 0.0395 |
| rc_AA426291_at | AA426291 | 961 | EST | 3.03 | 슠 | 0.00365 |
| rc_AA426374_f_at | AA426374 | 964 | tubulin, alpha 2 | 3.25 | d | 0.04346 |
| rc_AA426447_at | AA426447 | 965 | EST | 3.01 | g. | 0.02414 |
| rc_AA426521_at | AA426521 | 296 | Sjogren's syndrome nuclear autoantigen 1 | 3.33 | 롸 | 0.01163 |
| rc_AA427734_at | AA427734 | 977 | cholinergic receptor, nicotinic, epsilon polypeptide | 3.08 | đ | 0.04796 |
| AA428172_f_at | AA428172 | 986 | Notch (Drosophila) homolog 3 | 9.63 | 슠 | 0.00195 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------------|----------|--------|---|-------------|----------------|---------|
| | | | cofactor required for Sp1 transcriptional activation, | | | |
| rc AA428204 at | AA428204 | 286 | subunit 6 (77kD) | 3.08 | d _n | 0.00313 |
| rc_AA429470_at | AA429470 | 966 | EST | 3.2 | dn | 0.0153 |
| rc AA429472 at | AA429472 | 266 | DKFZP434P106 protein | 8.78 | d | 0.00063 |
| AA429539 f at | AA429539 | 666 | EST | 4.3 | ф | 0.01035 |
| rc AA429572 at | AA429572 | 1000 | ribosomal protein S6 | 3.31 | dn | 0.02144 |
| AA429825 at | AA429825 | 1003 | DKFZP566B023 protein | 3.11 | 슠 | 0.01857 |
| rc AA43003 <u>2</u> at | AA430032 | 1009 | pituitary tumor-transforming 1 | 10.67 | d | 0.00052 |
| rc_AA430048_at | AA430048 | 1012 | KIAA0160 protein | 4.32 | d | 0.00279 |
| rc_AA430154_at | AA430154 | 1014 | EST | 3.09 | 롸 | 0.04401 |
| rc_AA430474_at | AA430474 | 1015 | EST | 4.69 | g. | 0.00007 |
| rc_AA430675_at | AA430675 | 1019 | Fanconi anemia, complementation group G | 3.16 | dn | 0.01007 |
| rc_AA431571_at | AA431571 | 1024 | EST | 4.62 | g | 0.0174 |
| rc_AA431719_at | AA431719 | 1025 | EST | 3.19 | dn | 0.00294 |
| rc_AA433947_at | AA433947 | 1034 | EST | 3.09 | dn | 0.00253 |
| rc_AA434418_at | AA434418 | 1036 | KIAA1115 protein | 6.75 | dn | 0.0032 |
| rc AA435662 f at | AA435662 | 1039 | EST | 3.27 | dn | 0.0433 |
| rc AA435665 at | AA435665 | 1040 | EST | 3.94 | dn | 0.00274 |
| rc AA435681 s at | AA435681 | 1041 | EST | 3.07 | ф | 0.01166 |
| rc AA435748 at | AA435748 | 1044 | EST | 5.01 | dn | 0.01812 |
| rc aa435769 s at | AA435769 | 1046 | EST | 3.06 | dn | 0.00615 |
| AA442054 s at | AA442054 | 1067 | phospholipase C, gamma 1 (formerly subtype 148) | 4.94 | d | 0.04102 |
| rc AA442155 at | AA442155 | 1068 | transforming acidic coiled-coil containing protein 3 | 3.35 | dn | 0.00344 |
| AA442400 at | AA442400 | 1071 | hepatitis B virus x-interacting protein (9.6kD) | 3.02 | ф | 0.04037 |
| rc AA442763 at | AA442763 | 1072 | cyclin B2 | 3.49 | dn | 0.04176 |
| rc_AA443271_at | AA443271 | 1073 | KIAA0546 protein | 3.44 | dn | 0.00324 |
| rc AA443316 s at | AA443316 | 1075 | v-Ha-ras Harvey rat sarcoma viral oncogene homolog | 3.4 | dn | 0.00133 |
| rc AA443602 at | AA443602 | 1078 | EST | 5.71 | d | 0.00736 |
| rc_AA443802_at | AA443802 | 1081 | EST | 4.07 | g | 0.01546 |
| rc_AA446242_at | AA446242 | 1087 | EST | 6.3 | dn | 0.00169 |
| rc_AA446570_at | AA446570 | 1089 | EST | 3.12 | d | 0.02228 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| rc_AA446581_at | AA446581 | 1090 | DKFZP564P0462 protein | 4.04 | g | 0.00479 |
| rc_AA446970_at | AA446970 | 1098 | EST | 3.09 | g | 0.01627 |
| rc_AA447574_at | AA447574 | 1102 | EST | 4.12 | g | 0.00779 |
| rc_AA448252_at | AA448252 | 1114 | EST | ო | d | 0.00256 |
| rc_aa449073_s_at | AA449073 | 1117 | EST | 5.61 | 롸 | 0.01214 |
| rc_aa449431_s_at | AA449431 | 1124 | translation initiation factor IF2 | 3.76 | g. | 0.00571 |
| rc_AA449828_at | AA449828 | 1130 | EST | 3.35 | g | 0.01609 |
| rc_AA450247_at | AA450247 | 1133 | EST | 3.13 | g | 0.00531 |
| | | | hepatocellular carcinoma associated protein; breast | | • | |
| rc_AA451680_at | AA451680 | 1136 | cancer associated gene 1 | 3.85 | đ | 0.0018 |
| rc_AA451877_at | AA451877 | 1138 | EST | 4.6 | g. | 0.04045 |
| AA451992_at | AA451992 | 1140 | HSPC039 protein | 3.33 | g | 0.01696 |
| rc_AA452167_at | AA452167 | 1142 | EST | 3.29 | g | 0.03337 |
| AA452724_at | AA452724 | 1149 | programmed cell death 5 | 7.7 | dn | 0.00085 |
| rc_AA453628_at | AA453628 | 1154 | EST | 3.17 | dn | 0.00849 |
| rc_AA453656_at | AA453656 | 1155 | EST | 3.02 | d | 0.00958 |
| rc_AA453783_s_at | AA453783 | 1158 | EST | 4.07 | đ | 0.00786 |
| rc_AA454597_s_at | AA454597 | 1166 | EST | 4.23 | g | 0.00917 |
| rc_AA454830_at | AA454830 | 1170 | DKFZP586M2123 protein | 6.48 | dn | 0.00555 |
| AA454908_s_at | AA454908 | 1171 | KIAA0144 gene product | 6.39 | dn | 0.01835 |
| rc_AA455239_at | AA455239 | 1174 | chromosome-associated polypeptide C | 5.78 | đ | 0.00003 |
| rc_AA456415_at | AA456415 | 1192 | KIAA0537 gene product | 3.32 | dn | 0.00155 |
| rc_AA456583_s_at | AA456583 | 1193 | PL6 protein | 3.37 | dn | 0.00139 |
| rc_AA456646_at | AA456646 | 1196 | EST | 3.34 | dn | 0.0309 |
| rc_AA456852_at | AA456852 | 1199 | suppressor of white apricot homolog 2 | 3.66 | đ | 0.00614 |
| rc_AA458878_s_at | AA458878 | 1204 | EST | 5.49 | đ | 0.00977 |
| rc_AA458890_at | AA458890 | 1206 | EST | 3.27 | ф | 0.00079 |
| rc_AA459254_at | AA459254 | 1211 | EST | 6.22 | đ | 0.00001 |
| AA459542_s_at | AA459542 | 1218 | regulatory factor X-associated ankyrin-containing protein | 3.4 | đ | 0.00841 |
| rc_AA460665_at | AA460665 | 1230 | EST | 4.01 | dn · | 0.01866 |
| rc_aa460909_s_at | AA460909 | 1232 | EST | 5.02 | ф | 0.01354 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| rc AA461063 at | AA461063 | 1235 | EST | 4.3 | dn | 0.00074 |
| AA461282 s at | AA461282 | 1237 | dihydropyrimidinase-like 2 | 3.42 | dn | 0.02014 |
| rc AA461476 at | AA461476 | 1243 | EST | 3.72 | ф | 0.00744 |
| rc AA463254 s at | AA463254 | 1247 | histone deacetylase 3 | 4.01 | dn | 0.01856 |
| rc AA463934 at | AA463934 | 1253 | splicing factor 3b, subunit 4, 49kD | 3.15 | dn | 0.00952 |
| AA464043 s at | AA464043 | 1255 | EST | 3.99 | dn | 0.00056 |
| rc AA464251 at | AA464251 | 1257 | EST | 3.45 | dn | 0.02229 |
| rc AA464414 i at | AA464414 | 1258 | EST | 4.08 | dn | 0.02299 |
| rc AA464423 at | AA464423 | 1259 | EST | 3.06 | dn | 0.01416 |
| rc aa464722 s at | AA464722 | 1263 | DKFZP566C243 protein | 3.51 | dn | 0.00101 |
| rc AA464963 at | AA464963 | 1265 | EST | 4.77 | đ | 0.00086 |
| AA465000 s at | AA465000 | 1266 | EST | 3.86 | ф | 0.00431 |
| rc AA465093 at | AA465093 | 1267 | TIA1 cytotoxic granule-associated RNA-binding protein | 3.3 | ф | 0.01314 |
| rc_AA465218_at | AA465218 | 1268 | DKFZP586M1523 protein | 3.17 | dn | 0.00357 |
| rc_AA465342_at | AA465342 | 1271 | EST | 3.21 | dn | 0.01378 |
| rc_AA470156_at | AA470156 | 1276 | EST | 4.99 | dn | 0.0206 |
| AA471384 at | AA471384 | 1278 | divalent cation tolerant protein CUTA | 3.44 | 롸 | 0.01161 |
| rc AA476473 at | AA476473 | 1285 | EST | က | dn | 0.01324 |
| rc_AA476754_s_at | AA476754 | 1287 | EST | 3.18 | dn | 0.01696 |
| rc AA476944 at | AA476944 | 1288 | EST | 3.29 | ф | 0.00189 |
| rc_AA477316_at | AA477316 | 1290 | calumenin | 3.05 | dn | 0.00608 |
| rc AA477549 s at | AA477549 | 1291 | T-cell, immune regulator 1 | 4.84 | dn | 0.04096 |
| rc AA478017 at | AA478017 | 1295 | zyxin | 4.25 | dn | 0.01223 |
| rc_AA478300_at | AA478300 | 1298 | CD39-like 2 | 3.75 | dn | 0.00152 |
| rc_AA478415_at | AA478415 | 1299 | EST | 3.14 | dn | 0.0483 |
| rc_AA478422_at | AA478422 | 1301 | unc-51 (C. elegans)-like kinase 1 | 3.83 | dn | 0.00116 |
| rc AA478615 s at | AA478615 | 1305 | H1 histone family, member X | 3.09 | dn | 0.0499 |
| | | | disabled (Drosophila) homolog 2 (mitogen-responsive | | | |
| rc_AA478971_s_at | AA478971 | 1306 | phosphoprotein) | 3.25 | ф | 0.02698 |
| rc AA479096 at | AA479096 | 1308 | EST | 3.32 | ф | 0.00118 |
| rc_AA479139_s_at | AA479139 | 1310 | acid phosphatase 1, soluble | 3.42 | ф | 0.01853 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Sed ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|----------------|---------|
| rc_AA479881_at | AA479881 | 1317 | | 3.34 | 음 | 0.03289 |
| rc_AA481060_at | AA481060 | 1326 | EST | 3.08 | 음 | 0.00029 |
| rc_AA481420_at | AA481420 | 1327 | EST | 3.08 | g | 0.0206 |
| rc_AA482104_s_at | AA482104 | 1332 | non-metastatic cells 3, protein expressed in | 4.78 | g | 0.00135 |
| rc_AA482224_f_at | AA482224 | 1334 | putative type II membrane protein | 4.47 | dn | 0.0001 |
| AA482319_f_at | AA482319 | 1335 | putative type II membrane protein | 4.9 | 롸 | 0.00028 |
| AA482319_i_at | AA482319 | 1335 | putative type II membrane protein | 3.13 | d | 0.00071 |
| rc_AA485060_at | AA485060 | 1339 | EST | 3.83 | g | 0.03172 |
| rc_AA485084_s_at | AA485084 | 1340 | EST | 3.31 | g | 0.01232 |
| rc_AA485431_s_at | AA485431 | 1345 | EST | 3.81 | g | 0.00441 |
| rc_AA485697_at | AA485697 | 1346 | EST | 3.53 | dn | 0.03566 |
| rc_AA487218_at | AA487218 | 1355 | EST | 4.43 | g | 0.03198 |
| rc_AA487856_at | AA487856 | 1359 | KIAA0676 protein | 3.59 | dn | 0.01408 |
| rc_AA488074_at | AA488074 | 1360 | cell division cycle 42 (GTP-binding protein, 25kD) | 3.74 | dn | 0.01887 |
| rc_AA488432_at | AA488432 | 1361 | phosphoserine phosphatase | 4.2 | d | 0.00128 |
| rc_AA488872_s_at | AA488872 | 1363 | EST | 3.35 | g. | 0.03191 |
| rc_AA488892_at | AA48892 | 1364 | EST | 4.14 | <u>s</u> | 0.04766 |
| rc_AA489091_at | AA489091 | 1368 | EST | 3.58 | dn | 0.0002 |
| rc_AA489707_at | AA489707 | 1371 | EST | 3.5 | g | 0.03208 |
| rc_AA489712_at | AA489712 | 1372 | EST . | 4.69 | đ | 0.00587 |
| rc_AA490212_at | AA490212 | 1375 | H2A histone family, member Y | 3.52 | d n | 0.02202 |
| | | | solute carrier family 2 (facilitated glucose transporter), | | • | |
| AA491188_at | AA491188 | 1387 | member 3 | 5.04 | dn | 0.02291 |
| rc_AA491295_at | AA491295 | 1390 | calcium/calmodulin-dependent protein kinase kinase 2, | 3.71 | dn | 0.0103 |
| AA495857_at | AA495857 | 1394 | EST | 3.21 | g | 0.02243 |
| rc_AA496715_f_at | AA496715 | 1400 | spectrin SH3 domain binding protein 1 | 3.44 | d _n | 0.00069 |
| - | | | v-erb-b2 avian erythroblastic leukemia viral oncogene | | | |
| rc_AA496981_at | AA496981 | 1404 | homolog 3 | 5.82 | g | 0.00521 |
| rc_AA497018_at | AA497018 | 1406 | adenylate cyclase 1 (brain) | 4.81 | dn | 0.00352 |
| AA504413_at | AA504413 | 1413 | | 3.31 | dn | 0.00036 |
| rc_AA504512_s_at | AA504512 | 1415 | KIAA0943 protein | 5.72 | 롸 | 0.00384 |
| | | | | | | |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank Se | Sed ID Known Gene Name | Vame | Fold Change | Direction | Pvalue |
|------------------|-------------|---|-----------------------|-------------|-----------|---------|
| | | solute carrier fami | glucose transporter), | | | |
| rc AA505133 at | AA505133 14 | 1417 member 3 | | 12.21 | dn | 0.00169 |
| rc_AA505141_at | • | | | 3.08 | ф | 0.02327 |
| rc_AA521149_at | | | | 3.33 | dn | 0.00211 |
| rc_AA598405_at | | _ | RGS16 | 3.87 | 롸 | 0.00649 |
| rc_AA598447_at | | • | ceptor for tRNAs) | 3.5 | ф | 0.01201 |
| rc_AA598589_at | | | | 3.24 | ф | 0.00432 |
| 1 | | SWI/SNF related, matrix associated, actin dependent | ted, actin dependent | | | |
| rc AA598648 s at | AA598648 | 1432 regulator of chromatin, subfamily a, member 4 | a, member 4 | 3.46 | 음 | 0.00293 |
| rc AA598712 at | | | | 3.45 | ď | 0.00005 |
| rc AA598749 at | | 1438 EST | | 3.01 | 슠 | 0.03714 |
| | | heterogeneous nuclear ribonucleoprotein U (scaffold | oprotein U (scaffold | | | |
| rc AA598829 s at | AA598829 | 1439 attachment factor A) | | 3.04 | ф | 0.00967 |
| rc_AA598831 f at | AA598831 | 1440 EST | | 3.41 | ф | 0.00452 |
| rc AA599469 at | | 1450 EST | | 3.07 | ᅀ | 0.04154 |
| rc_AA599808_at | | 1455 EST | | 3.09 | dn | 0.00726 |
| rc_AA599850_at | | | | 3.55 | dn | 0.03215 |
| rc_AA600153_at | | | | 3.71 | dn | 0.02967 |
| rc_AA608668_at | | _ | and 4.1-like 2 | 3.33 | dn | 0.02014 |
| rc_AA608897_at | AA608897 | | | 5.05 | d | 0.01782 |
| rc_AA609008_at | • | | | 4.04 | dn | 0.00002 |
| rc_AA609080_at | | 1478 EST | | 3.71 | d | 0.0306 |
| rc_AA610073_at | • | 1497 EST | | 3.25 | dn | 0.00859 |
| rc_AA610089_at | • | 1498 U4/U6-associated RNA splicing factor | factor | 4.07 | ф | 0.00361 |
| | • | 1499 tetraspan NET-6 protein | | 16.35 | d | 0.00249 |
| rc AA620461 at | • | | | 3.45 | d | 0.01146 |
| rc AA620553 s at | _ | | ase 1 | 7.56 | dn | 0.00101 |
| rc AA620761 at | | | | 3.3 | dn | 0.00285 |
| rc_AA620779_at | _ | 508 golgin-67 | | 3.35 | dn | 0.00297 |
| rc AA620881 at | • | | | 9.49 | dn | 0.00062 |
| rc_AA621146_at | AA621146 | | | 3.15 | ф | 0.02116 |
| 1 | | | | | | |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|--|-------------|-----------|---------|
| rc_AA621242_s_at | AA621242 | 1518 | hypothetical protein, peptidylprolyl isomerase B | 4.59 | 롸 | 0.00081 |
| rc_AA621367_at | AA621367 | 1523 | EST | 3.1 | 9 | 0.00066 |
| rc_AA621409_at | AA621409 | 1524 | putative type II membrane protein | 3.5 | d | 0.00462 |
| rc_AA621530_at | AA621530 | 1526 | EST | 3.26 | g | 0.00298 |
| rc_AA621535_at | AA621535 | 1527 | FE65-LIKE 2 | 3.37 | dn | 0.0167 |
| rc_AA621752_at | AA621752 | 1529 | 26S proteasome-associated pad1 homolog | 3.13 | dn | 0.01571 |
| AB002373_at | AB002373 | 1538 | KIAA0375 gene product | 4.41 | d | 0.00795 |
| AF003521_at | AF003521 | 1545 | jagged 2 | 3.58 | đ | 0.00299 |
| AF004022_at | AF004022 | 1546 | serine/threonine kinase 12 | 3.29 | d | 0.00841 |
| C00358_at | C00358 | 1552 | nucleolar protein 3 (apoptosis repressor with CARD | 3.45 | dn | 0.00985 |
| C01721_at | C01721 | 1558 | phospholipase C, beta 3, neighbor pseudogene | 5.89 | dn | 0.0383 |
| C01766_s_at | C01766 | 1559 | EST | 8.18 | dn | 0.00505 |
| rc_C14051_f_at | C14051 | 1565 | phosphoprotein enriched in astrocytes 15 | 4.79 | 9 | 0.00548 |
| rc_C14098_f_at | C14098 | 1566 | EST | 4.62 | d | 0.01654 |
| rc_C14756_f_at | C14756 | 1570 | MLN51 protein | 3.75 | d | 0.0226 |
| rc_C14835_f_at | C14835 | 1571 | EST | 3.35 | d | 0.0316 |
| D00596_at | D00596 | 1590 | thymidylate synthetase | 5.58 | đ | 0.0098 |
| D13370_at | D13370 | 1603 | APEX nuclease (multifunctional DNA repair enzyme) | 3.07 | g | 0.00857 |
| | | | general transcription factor IIIC, polypeptide 2 (beta | | | |
| D13636_at | D13636 | 1606 | subunit, 110kD) | 3.12 | d | 0.00022 |
| D13640_at | D13640 | 1608 | KIAA0015 gene product | 3.55 | dn | 0.00347 |
| D14657_at | D14657 | 1615 | KIAA0101 gene product | 3.84 | dn | 0.02048 |
| rc_D20899_at | D20899 | 1626 | EST | 3.13 | dn | 0.02128 |
| | | | minichromosome maintenance deficient (S. cerevisiae) 2 | 21 | | |
| D21063_at | D21063 | 1628 | (mitotin) | 3.25 | 음 | 0.03558 |
| D26129_at | D26129 | 1635 | ribonuclease, RNase A family, 1 (pancreatic) | 6.9 | g | 0.00008 |
| D28589_at | D28589 | 1637 | EST | 3.38 | dn | 0.01144 |
| D30946_at | D30946 | 1638 | kinesin family member 3B | 3.43 | dn | 0.01458 |
| D31094_at | D31094 | 1639 | G8 protein | 9.37 | dn | 0.0048 |
| D31294_at | D31294 | 1643 | EST | 3.3 | dn | 0.004 |
| D31417_at | D31417 | 1645 | secreted protein of unknown function | 3.69 | ф | 0.0004 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|--|-------------|------------|---------|
| D38073 at | D38073 | 1651 | minichromosome maintenance deficient (S. cerevisiae) 3 | 4.1 | ф | 0.01195 |
| D38305_at | D38305 | 1652 | transducer of ERBB2, 1 | 3.22 | dn | 0.0215 |
| D42040 s at | D42040 | 1657 | female sterile homeotic-related gene 1 (mouse homolog) | 4.02 | dn | 0.00389 |
| rc D51072 s at | D51072 | 1674 | biliverdin reductase A | 3.34 | dn | 0.0254 |
| rc_D51276_f at | D51276 | 1678 | leukemia-associated phosphoprotein p18 (stathmin) | 9.42 | dn | 0.00015 |
| D55716 at | D55716 | 1686 | minichromosome maintenance deficient (S. cerevisiae) 7 | 5.48 | dn | 0.00003 |
| rc D57317 at | D57317 | 1688 | activated RNA polymerase II transcription cofactor 4 | 3.17 | ф | 0.00464 |
| rc D59355 s at | D59355 | 1696 | cytoskeleton-associated protein 1 | 6.05 | ф | 0.0015 |
| rc_D59553_f_at | D59553 | 1697 | golgin-67 | 5.95 | dn | 0.00169 |
| rc_D59570_f at | D59570 | 1699 | EST | 4.34 | dn | 0.00487 |
| rc_D60811_s_at | D60811 | 1704 | EST | 4.34 | dn | 0.00217 |
| D63478 at | D63478 | 1711 | KIAA0144 gene product | 3.89 | ф | 0.00253 |
| D63486_at | D63486 | 1712 | KIAA0152 gene product | 3.56 | ф | 0.00063 |
| rc D80420 at | D80420 | 1732 | ubiquinol-cytochrome c reductase hinge protein | 3.86 | đ | 0.00412 |
| rc D80710 f at | D80710 | 1734 | integral type I protein | 3.17 | dn | 0.04549 |
| rc D80917 f at | D80917 | 1736 | KIAA0670 protein/acinus | 3.09 | dn | 0.00168 |
| rc_D80946_f_at | D80946 | 1737 | SFRS protein kinase 1 | 3.07 | dn | 0.00986 |
| | D81608 | 1740 | polymerase (RNA) II (DNA directed) polypeptide K | 3.52 | dn | 0.00437 |
| D82226 s at | D82226 | 1742 | proteasome (prosome, macropain) 26S subunit, | 4.35 | đ | 0.00184 |
| D82277 s at | D82277 | 1743 | LDL induced EC protein | 3.33 | ф | 0.00355 |
| D82558 at | D82558 | 1746 | novel centrosomal protein RanBPM | 4.67 | d | 0.00458 |
| ! | | | trinucleotide repeat containing 11 (THR-associated | | | |
| D83783_at | D83783 | 1748 | protein, 230 kDa subunit) | 4.16 | d | 0.00055 |
| D84557_at | D84557 | 1749 | minichromosome maintenance deficient (mis5, S. | 3.97 | 읔 | 0.0017 |
| D86957_at | D86957 | 1754 | KIAA0202 protein | 3.08 | dn | 0.02949 |
| D86977 at | D86977 | 1757 | KIAA0224 gene product | 3.03 | dn | 0.00053 |
| rc_F01538_s_at | F01538 | 1771 | RAP1, GTPase activating protein 1 | 4.88 | 슠 | 0.00292 |
| rc F01568 at | F01568 | 1772 | EST | 4.13 | 롸 | 0.00084 |
| rc F01831 at | F01831 | 1773 | EST | 5.95 | ф | 0.00532 |
| rc F02254 s at | F02254 | 1777 | Fas-activated serine/threonine kinase | 5.1 | g | 0.00329 |
| rc_F02807_at | F02807 | 1781 | KIAA0838 protein | 2.67 | <u>다</u> : | 0.02064 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|-----------------|---------|--------|---|-------------|-----------|---------|
| rc_F02863_at | F02863 | 1782 | EST | 3.05 | dn | 0.03504 |
| rc_F04320_s_at | F04320 | 1786 | replication factor C (activator 1) 4 (37kD) | 6.29 | dn | 0.00042 |
| rc_F04444_at | F04444 | 1788 | EST | 4.13 | dn | 0.00944 |
| rc_F04479_at | F04479 | 1789 | KIAA1067 protein | 3.23 | dn | 0.04522 |
| rc_F08876_at | F08876 | 1797 | EST | 90.6 | d | 0 |
| | | | procollagen-proline, 2-oxoglutarate 4-dioxygenase | | | |
| rc_F09788_at | F09788 | 1808 | (proline 4-hydroxylase), alpha polypeptide II | 3.67 | d | 0.01682 |
| rc_F10199_f_at | F10199 | 1813 | EST . | 3.93 | 由 | 0.03209 |
| rc_F10290_at | F10290 | 1815 | EST | 3.39 | dn | 0.02392 |
| rc_F10453_at | F10453 | 1819 | EST | 3.64 | dn | 0.01878 |
| rc_F10741_at | F10741 | 1822 | KIAA0622 protein | 3.01 | d | 0.03079 |
| rc_F13809_f_at | F13809 | 1828 | tropomyosin 1 (alpha) | 4.4 | ď | 0.01221 |
| rc_H00540_at | H00540 | 1829 | EST | 3.74 | đ | 0.00234 |
| rc_H05084_at | H05084 | 1844 | EST | 5.85 | d | 0.0059 |
| rc_H07873_at | H07873 | 1856 | EST | 3.53 | dn | 0.0391 |
| rc_H08863_at | H08863 | 1859 | hypothetical protein | 7.18 | d | 0.02102 |
| rc_H09241_s_at | H09241 | 1861 | EST | 3.05 | đ | 0.01487 |
| rc_H09271_f_at | H09271 | 1862 | EST | 4.78 | d | 0.00072 |
| rc_H10933_at | H10933 | 1873 | EST | 6.18 | dn | 0.00003 |
| rc_H11320_s_at | H11320 | 1875 | SUMO-1 activating enzyme subunit 2 | 3.06 | d | 0.00167 |
| rc_H16251_s_at. | H16251 | 1886 | EST | 3.3 | g | 0.03286 |
| rc_H27188_f_at | H27188 | 1908 | collagen-binding protein 2 (colligen 2) | 5.84 | dn | 0.01826 |
| rc_H27897_s_at | H27897 | 1911 | hypothetical protein | 3.01 | g | 0.00174 |
| rc_H28333_f_at | H28333 | 1912 | melanoma adhesion molecule | 4.94 | dn | 0.00166 |
| rc_H41529_at | H41529 | 1926 | EST | 5.06 | dn | 0.03309 |
| H46486_s_at | H46486 | 1932 | nesca protein | 4.57 | d | 0.00749 |
| rc_H47357_f_at | H47357 | 1934 | EST | 3.65 | đ | 0.03799 |
| rc_H48459_s_at | H48459 | 1937 | KIAA0186 gene product | 3.1 | ф | 0.02325 |
| rc_H52937_at | H52937 | 1944 | roundabout (axon guidance receptor, Drosophila) | 4.02 | dn | 0.00163 |
| rc_H56345_r_at | H56345 | 1950 | EST | 3.73 | ф | 0.00853 |
| rc_H57709_s_at | H57709 | 1956 | ribosomal protein L31 | 4.41 | ф | 0.00091 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| rc H59617 at | H59617 | 1964 | EST | 5.81 | ф | 0.0115 |
| rc H62474 f at | H62474 | 1970 | EST | 3.39 | dn | 0.04173 |
| rc_H64493_f_at | H64493 | 1973 | immunoglobulin heavy constant gamma 3 (G3m marker) | 4.74 | ф | 0.00751 |
| | | | phospholipase A2, group VII (platelet-activating factor | | | |
| rc H65030 s at | H65030 | 1974 | acetylhydrolase, plasma) | 3.26 | dn | 0.02278 |
| rc H65042 at | H65042 | 1975 | EST | 3.44 | ф | 90000 |
| H67964 at | H67964 | 1981 | EST | 3.06 | ᅀ | 0.02707 |
| rc H68794 at | H68794 | 1984 | EST | 3.67 | dn | 0.00327 |
| rc H70739 f at | H70739 | 1991 | EST | 4.34 | dn | 0.00106 |
| rc_H73484_s_at | H73484 | 1995 | ferritin, heavy polypeptide 1 | 3.18 | dn | 0.00432 |
| rc H78211 at | H78211 | 2001 | EST | 7.5 | ф | 0.02674 |
| rc H86072 f at | H86072 | 2015 | EST | 4.49 | ф | 0.01301 |
| rc H88674 s at | H88674 | 2021 | collagen, type I, alpha 2 | 4.15 | dn | 0.02664 |
| rc_H89987_s_at | H89987 | 2027 | ATP-binding cassette, sub-family C (CFTR/MRP), | 3.13 | dn | 0.01194 |
| rc H91632 at | H91632 | 2031 | EST | 3.5 | dn | 0.03688 |
| rc_H94471_at | H94471 | 2042 | occludin | 6.26 | dn | 0.00379 |
| rc_H96850_at | H96850 | 2055 | dolichyl-diphosphooligosaccharide-protein | 3.03 | g | 0.00679 |
| rc_H97012_at | H97012 | 2058 | EST | 3.51 | dn | 0.03505 |
| rc_H97013_at | H97013 | 2059 | ephrin-A4 | 8.9 | 롸 | 0.00023 |
| rc H97677 s at | H97677 | 2062 | EST | 4.34 | ф | 0.00753 |
| rc_H99261 s at | H99261 | 2074 | EST | 3.33 | ф | 0.00319 |
| rc H99364 at | H99364 | 2075 | chloride channel 7 | 3.03 | dn | 0.01727 |
| rc H99473 s at | H99473 | 2077 | regulator of nonsense transcripts 1 | 6.51 | dn | 0.00025 |
| rc_H99489 s_at | H99489 | 2078 | quiescin Q6 | 3.4 | dn | 0.02682 |
| rc_H99587_s_at | H99587 | 2079 | EST | 4.44 | dn | 0.00532 |
| rc H99774 at | H99774 | 2081 | EST | 3.51 | dn | 60000.0 |
| rc_H99877_at | H99877 | 2083 | exportin, tRNA (nuclear export receptor for tRNAs) | 3.75 | d | 0.00302 |
| rc_H99879_at | H99879 | 2084 | EST | 10.81 | d | 0.001 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| J00231 f at J00231 J03040_at J03040 J03464_s_at J03464 J04029_s_at J04029 J05614_at J05614 L03411_s_at L03411 L04270_at L04270 L06797_s_at L06797 L11669_at L17131 L25444_at L25444 L25876_at L25876 L25876_at L25876 L29218_s_at L29218 L29218_s_at L29218 | 2087 | | | | |
|---|------|---|-------|--------------|---------|
| · | 2087 | | , | | |
| · | | immunoglobulin heavy constant gamma 3 (G3m marker) | 6.23 | dn | 0.00177 |
| • | 2091 | secreted protein, acidic, cysteine-rich (osteonectin) | 3.77 | dn | 0.00594 |
| · | 2094 | collagen, type I, alpha 2 | 10.37 | dn | 0.00979 |
| | | keratin 10 (epidermolytic hyperkeratosis; keratosis | | | |
| | 2102 | palmaris et plantaris) | 3.02 | dn | 0.00032 |
| | 2122 | EST | 3.73 | on | 0.03419 |
| | 2134 | RD RNA-binding protein | 4.06 | g n | 0.00467 |
| | 2135 | lymphotoxin beta receptor (TNFR superfamily, member | 3.5 | <u>a</u> | 0.01547 |
| | 2143 | chemokine (C-X-C motif), receptor 4 (fusin) | 3.23 | g n | 0.04782 |
| | 2157 | tetracycline transporter-like protein | 3.4 | g | 0.02062 |
| | | high-mobility group (nonhistone chromosomal) protein | | | |
| | 2168 | isoforms I and Y | 4.45 | an | 0.03141 |
| | ٠ | TATA box binding protein (TBP)-associated factor, RNA | | • | |
| | 2181 | polymerase II, E, 70/85kD | 3.78 | dn | 0.00011 |
| | | cyclin-dependent kinase inhibitor 3 (CDK2-associated | | | |
| | 2182 | dual specificity phosphatase) | 4.43 | dn | 0.00082 |
| | 2188 | mannosidase, alpha, class 2A, member 2 | 4.63 | dn | 0.00876 |
| | 2190 | CDC-like kinase 2 | 6.51 | dn | 0.00019 |
| | 2190 | CDC-like kinase 2 | 3.82 | ď | 0.00035 |
| L33930_s_at L33930 | 2198 | CD24 antigen (small cell lung carcinoma cluster 4 | 4.35 | g. | 0.03968 |
| | | transcription elongation factor B (SIII), polypeptide 1 | | | |
| L34587_at L34587 | 2200 | (15kD, elongin C) | 3.12 | dn | 0.00946 |
| L47125_s_at L47125 | 2218 | glypican 3 | 10.69 | 9 | 0.04129 |
| L76191_at L76191 | 2222 | interleukin-1 receptor-associated kinase 1 | 3.85 | dn | 0.00152 |
| 1 76568 xnt3 f at 1 76568 | 2225 | deficiency complementation droup 4 | " | | 0.0172 |
| . < | 2241 | tronomyosin 2 (heta) | 7 13 | } | 0.00 |
| , de | 2261 | prothymosin, alpha (gene seguence 28) | 3.36 | } | 0.00033 |
| 1 | 2286 | tropomyosin 1 (alpha) | 4.3 | - | 0.00893 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seg ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|----------|--------|---|-------------|-----------|---------|
| M21259 at | M21259 | 2293 | small nuclear ribonucleoprotein polypeptide E | 3.68 | ф | 0.00415 |
| M26576 cds2 at | M26576 | 2310 | EST | 3.48 | dn | 0.00062 |
| AFFX-M27830 5 at | M27830 | 2314 | EST | 4.6 | dn | 0.04719 |
| AFFX-M27830 M at | M27830 | 2314 | EST | 3.54 | 습 | 0.00777 |
| AFFX-M27830 5 at | M27830 | 2314 | EST | 3.3 | ф | 0.02453 |
| | M31303 | 2327 | leukemia-associated phosphoprotein p18 (stathmin) | 5.86 | dn | 0.00071 |
| M32977 s at | M32977 | 2336 | vascular endothelial growth factor | 3.93 | dn | 0.04917 |
| - | | | v-erb-b2 avian erythroblastic leukemia viral oncogene | | | |
| M34309 at | M34309 | 2342 | homolog 3 | 3.49 | d | 0.00191 |
| M35252 at | M35252 | 2343 | transmembrane 4 superfamily member 3 | 4.65 | dn | 0.04128 |
| M37583 at | M37583 | 2349 | H2A histone family, member Z | 4.25 | dn | 0.00135 |
| M55210 at | M55210 | 2353 | laminin, gamma 1 (formerly LAMB2) | 3.47 | dn | 0.02551 |
| M55998 s at | M55998 | 2356 | collagen, type I, alpha 1 | 3.54 | dn | 0.01449 |
| | . M57710 | 2357 | lectin, galactoside-binding, soluble, 3 (galectin 3) | 6.76 | dn | 0.00103 |
| M57730 at | M57730 | 2358 | ephrin-A1 | 3.39 | dn | 0.00199 |
| M60784 s at | M60784 | 2366 | small nuclear ribonucleoprotein polypeptide A | 4.74 | ф | 0.00001 |
| M61916 at | M61916 | 2372 | laminin, beta 1 | 3.18 | ф | 0.01171 |
| M63573_at | M63573 | 2377 | peptidylprolyl isomerase B (cyclophilin B) | 3.59 | d | 0.00916 |
| M68864 at | M68864 | 2389 | ORF | 3.95 | dn | 0.00144 |
| M86667 at | M86667 | 2411 | nucleosome assembly protein 1-like 1 | 3.08 | d | 0.00473 |
| | | | stress-induced-phosphoprotein 1 (Hsp70/Hsp90- | | | |
| M86752 at | M86752 | 2412 | organizing protein) | 5.15 | g | 0.02881 |
| M87339_at | M87339 | 2415 | replication factor C (activator 1) 4 (37kD) | 4.59 | d | 0.00116 |
| M91083 at | M91083 | 2419 | chromosome 11 open reading frame 13 | 3.19 | dn | 0.00243 |
| l | | | membrane component, chromosomal 4, surface marker | | | |
| M93036 at | M93036 | 2422 | (35kD glycoprotein) | 3.07 | dn | 0.04199 |
| M94250_at | M94250 | 2426 | midkine (neurite growth-promoting factor 2) | 9.86 | dn | 0.02104 |
| M94345 at | M94345 | 2427 | capping protein (actin filament), gelsolin-like | 3.59 | ᅀ | 0.04508 |
| M97856 at | M97856 | 2436 | nuclear autoantigenic sperm protein (histone-binding) | 3.21 | 음 | 0.00444 |
| rc N21407 at | N21407 | 2443 | EST | 3.47 | 함 | 0.01037 |
| rc_N21648_s_at | N21648 | 2447 | MpV17 transgene, murine homolog, glomerulosclerosis | 3.73 | ф | 0.00071 |
| | | | | | | |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|-----------|---------|
| rc_N26904_at | N26904 | 2469 | EST | 15.38 | 롸 | 0.00077 |
| rc_N29484_at | N29484 | 2478 | EST | 3.08 | dn | 0.04834 |
| rc_N29742_at | N29742 | 2481 | EST | 3.74 | dn | 0.00104 |
| rc_N31597_s_at | N31597 | 2487 | DKFZP564G2022 protein | 3.17 | d | 0.03017 |
| rc_N33920_at | N33920 | 2493 | diubiquitin | 50.29 | d | 0 |
| rc_N34825_s_at | N34825 | 2498 | DKFZP434P106 protein | 3.27 | dn | 0.01334 |
| rc_N35913_at | N35913 | 2503 | EST | 3.48 | dn | 0.0016 |
| N36432_at | N36432 | 2507 | erythrocyte membrane protein band 4.1-like 2 | 7.95 | dn | 0.00067 |
| rc_N39237_at | N39237 | 2511 | EST | 3.45 | dn | 0.02481 |
| N42272_s_at | N42272 | 2515 | EST | 3.03 | dn | 0.0017 |
| | | | eukaryotic translation initiation factor 3, subunit 3 | | | |
| rc_N47956_at | N47956 | 2524 | (gamma, 40kD) | 3.76 | 음 | 0.00968 |
| rc_N48790_at | N48790 | 2532 | EST | 3.32 | dn | 0.00654 |
| rc_N51590_s_at | N51590 | 2546 | EST | 3.01 | dn | 0.04345 |
| rc_N51771_at | N51771 | 2548 | KIAA0652 gene product | 3.5 | dn | 0.00028 |
| | | | ADP-ribosyltransferase (NAD+; poly (ADP-ribose) | | | |
| rc_N51855_at | N51855 | 2550 | polymerase)-like 3 | 3.39 | đ | 0.00115 |
| rc_N52168_at | N52168 | 2551 | EST | 3.66 | dn | 0.00127 |
| rc_N53067_at | N53067 | 2557 | DKFZP547E1010 protein | 3.1 | dn | 0.00101 |
| rc_N54067_at | N54067 | 2562 | mitogen-activated protein kinase kinase kinase 4 | 4.82 | 롸 | 0.00229 |
| rc_N54841_at | N54841 | 2572 | EST | 5.87 | dn | 0.02752 |
| rc_N56935_s_at | N56935 | 2575 | EST | 4.04 | 슠 | 0.00797 |
| rc_N59536_at | N59536 | 2586 | EST | 11.68 | dn | 0.00484 |
| rc_N62126_at | N62126 | 2589 | EST | 6.42 | 요 | 0.00109 |
| rc_N64374_at | N64374 | 2608 | KIAA0537 gene product | 3.25 | ф | 0.01652 |
| rc_N67815_f_at | N67815 | 2627 | EST | 3.84 | dn | 0.00439 |
| rc_N68018_at | N68018 | 2631 | TBP-associated factor 172 | 3.84 | dn | 0.00277 |
| rc_N68241_at | N68241 | 2634 | EST | 4.32 | d d | 0.00532 |
| rc_N69084_i_at | N69084 | 2643 | EST | 3.11 | dn | 0.0094 |
| rc_N69252_f_at | N69252 | 2647 | ferritin, light polypeptide | 3.69 | g, | 0.04116 |
| rc_N69263_at | N69263 | 2648 | EST | 5.26 | 롸 | 0.0276 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genhank Sed ID | ID Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|----------------|--|-------------|-----------|---------|
| rc N69390 at | | EST | 3.99 | g | 0.00016 |
| rc N69879 s at | | | 3,15 | dn | 0.01659 |
| rc N70481 at | | 8 EST | 4.13 | dn | 0.0099 |
| rc N70678 s at | | | 3.78 | 롸 | 0.02858 |
| 1 | | solute carrier family 11 (proton-coupled divalent metal | | | |
| rc N72116 s at | N72116 2668 | | 5.57 | dn | 0.00709 |
| rc N73705 at | N73705 2677 | _ | 3.75 | ᅀ | 0.01762 |
| | | | 3.67 | 음 | 0.00796 |
| rc N73808 f at | | | 6.44 | dn | 0.00352 |
| · | | | 4.43 | ᅀ | 0.00177 |
| rc_N75541_at | | | 4.43 | 롸 | 0.01059 |
| rc_N80703_at | | 4 EST | 5.65 | 롸 | 0.0001 |
| rc_N90238 i at | N90238 2712 | 2 EST | 3.13 | 롸 | 0.02492 |
| rc N91773 at | N91773 2719 | | 4.31 | ф | 0.00302 |
| rc N92948 s at | N92948 2726 | 6 nuclear phosphoprotein similar to S. cerevisiae PWP1 | 4.09 | 습 | 0.0019 |
| rc_N93299 f at | N93299 2732 | 2 nuclear receptor co-repressor 1 | 6.99 | 함 | 0.0371 |
| rc N93316 at | | | 3.16 | 롸 | 0.01262 |
| rc_N93798_at | N93798 2738 | 18 protein tyrosine phosphatase type IVA, member 3 | 4.91 | 롸 | 0.00245 |
| rc N98464 s at | N98464 2744 | | 3.68 | ф | 0.03007 |
| rc_N98758 f at | N98758 2745 | 5 EST | 3.54 | ф | 0.02609 |
| ¦ဖ | N99944 2749 | 9 EST | 3.46 | dn | 0.00104 |
| | R05316 2760 | 0 EST | 4.2 | dn | 0.00011 |
| rc_R06251_f_at | R06251 27 | 2764 tumor protein D52-like 2 | 4.88 | dn | 0.03097 |
| rc_R06254 f at | R06254 27 | 2765 tumor protein D52-like 2 | 3.53 | dn | 0.04865 |
| rc R06400 at | R06400 27 | 2768 EST | 3.03 | d | 0.03266 |
| rc R06986 f at | R06986 27 | 2776 peptidylprolyl isomerase B (cyclophilin B) | 7.03 | dn | 0.00628 |
| rc_R07172 i at | R07172 2777 | | | đ | 0.01322 |
| rc R15740 at | R15740 2 | 2791 carbohydrate (keratan sulfate Gal-6) sulfotransferase 1 | | ဌ | 0.00268 |
| rc R16144 at | R16144 2 | | | d | 0.0087 |
| rc_R20817_s_at | R20817 2 | | 3.03 | 앜 | 0.01091 |
| rc_R22565_at | R22565 28 | 2800 EST | 3.52 | 珨 | 0.04352 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank Sed ID | C Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|----------------|---|-------------|-----------|---------|
| rc_R26744_at | | midline 1 (Opitz/BBB syndrome) | 3.8 | 유 | 0.00266 |
| l | | myristoylated alanine-rich protein kinase C substrate | | | |
| rc R27016 s at | R27016 2806 | (MARCKS, 80K-L) | 3.53 | d | 0.03056 |
| rc R27296 f at | | ш | 3.41 | d n | 0.00309 |
| | | UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, | - | | |
| rc_R28636_at | R28636 2809 | | 3.06 | g S | 0.03678 |
| rc_R31607_at | R31607 2813 | EST | 3.41 | ဌ | 0.00163 |
| rc_R33498_s_at | R33498 2820 | EST | 3.5 | dn | 0.03336 |
| rc R39191 s at | R39191 2834 | KIAA1020 protein | 5.18 | g | 0.03185 |
| R39390_at | R39390 2837 | EST | 4.18 | dn | 0.0004 |
| rc_R39610_s_at | R39610 2838 | calpain, large polypeptide L2 | 3.13 | g | 0.01863 |
| rc_R43952_at | R43952 2853 | homeo box B5 | 3.39 | dn | 0.04829 |
| rc_R44617 f at | R44617 2857 | MyoD family inhibitor | 6.54 | dn | 0.02505 |
| rc_R44793_at | R44793 2859 | EST | 5.4 | dn | 0.00329 |
| rc_R44839_at | R44839 2861 | i-beta-1,3-N-acetylglucosaminyltransferase | 2 | dn | 0.01812 |
| rc_R45569_at | R45569 2864 | DKFZP547E1010 protein | 3.96 | dn | 0.00259 |
| rc_R45994_f_at | R45994 2867 | EST | 6.48 | d n | 0.00358 |
| rc_R46079_f_at | R46079 2868 | EST | 3.03 | dn | 0.00755 |
| rc_R46337_s_at | R46337 2869 | secretory carrier membrane protein 3 | 3.01 | dn | 0.00374 |
| rc_R48447_at | R48447 2871 | EST | 4.76 | dn | 0.00533 |
| rc_R48473_f_at | R48473 2872 | _ | 3.46 | dn | 0.01196 |
| rc_R48594_s_at | R48594 2875 | | 6.15 | dn | 0.03831 |
| rc_R49395_s_at | R49395 2881 | EST | 3.31 | dn | 0.00867 |
| rc_R49476_at | R49476 2883 | _ | 4.93 | 合 | 0.00763 |
| rc_R49482_at | R49482 2884 | EST | 3.27 | dn | 0.0161 |
| rc_R49708_s_at | R49708 2886 | EST | 4.56 | đ | 0.03767 |
| rc_R51908_s_at | R51908 2892 | | 3.16 | dn | 0.0083 |
| rc_R52161_at | R52161 2893 | EST | 3.41 | ф | 0.00053 |
| rc_R52649_at | R52649 2894 | EST | 4.69 | ф | 0.00135 |
| rc_R53109_f_at | R53109 2899 | | 3.31 | dn | 0.02406 |
| rc_R54614_s_at | R54614 2902 | EST | 3.22 | dn | 0.00334 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affv ID | Genbank | Sea ID | Known Gene Name | Fold Change | Direction | Pvalue |
|-----------------|---------|--------|---|-------------|-----------|---------|
| rc R56095 s at | R56095 | 2906 | EST | 3.67 | dn | 0.0158 |
| rc_R60512 s at | R60512 | 2918 | KIAA0191 protein | 3.08 | d | 0.00856 |
| rc R61374 at | R61374 | 2921 | EST | 4.33 | 롸 | 0.01489 |
| rc_R61557_at | R61557 | 2922 | KIAA0100 gene product | 3.9 | ф | 0.00292 |
| rc_R62456_at | R62456 | 2925 | _ | 3.44 | dn | 0.00285 |
| rc _R66469 f at | R66469 | 2937 | pleckstrin and Sec7 domain protein | 3.52 | dn | 0.0272 |
| rc R70005 at | R70005 | 2944 | EST | 4.98 | dn | 0.00007 |
| rc_R70253_at | R70253 | 2945 | EST | 3.38 | dn | 0.03125 |
| rc_R70532_at | R70532 | 2947 | EST | 3.44 | dn | 0.02186 |
| rc R70801 s at | R70801 | 2950 | EST | 90'9 | dn | 0.00291 |
| rc R71395 at | R71395 | 2952 | EST | 4.12 | dn | 0.03719 |
| rc R72886 s at | R72886 | 2956 | KIAA0422 protein | 5.5 | dn | 0.00091 |
| rc R73569 s at | R73569 | 2960 | EST | 3.54 | d | 0.01962 |
| } | | | O-linked N-acetylglucosamine (GlcNAc) transferase | | | |
| | | | (UDP-N-acetylglucosamine:polypeptide-N- | | | |
| rc R76782 s at | R76782 | 2963 | acetylglucosaminyl transferase) | 3.73 | dn | 0.00094 |
| rc_R77451_i_at | R77451 | 2964 | | 3.67 | ď | 0.00078 |
| rc_R79246_f_at | R79246 | 2969 | melanoma adhesion molecule | 90.9 | dn | 0.00057 |
| rc R91753 at | R91753 | 2983 | EST | 3.45 | ф | 0.02391 |
| rc_R91819_at | R91819 | 2984 | EST | 12.81 | dn | 0.00037 |
| rc R92449 s at | R92449 | 2985 | KIAA0323 protein | 4.34 | dn | 0.00104 |
| rc_R96527_s_at | R96527 | 2999 | KIAA0253 protein | 4.62 | dn | 0.00702 |
| rc_R96924_s_at | R96924 | 3001 | _ | 7.04 | ф | 0.00012 |
| S67070 at | S67070 | 3023 | heat shock 27kD protein 2 | 3.12 | đ | 0.01688 |
| S78187 at | S78187 | 3036 | cell division cycle 25B | 4.83 | d | 0.00547 |
| rc T03438 s at | T03438 | 3043 | EST | 3.79 | d | 0.02042 |
| rc_T03580_f at | T03580 | 3046 | pyruvate kinase, muscle | 5.57 | dn | 0.01344 |
| rc T03749 at | T03749 | 3048 | KIAA1089 protein | 4.23 | 由 | 0.00776 |
| rc T10316 s at | T10316 | 3052 | EST | 3.2 | 롸 | 0.04794 |
| rc_T10698_s_at | T10698 | 3054 | | 3.86 | 롸 | 0.00195 |
| rc_T15852_f_at | T15852 | 3062 | EST | 5.21 | ф | 0.00642 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank Se | Seq ID Known Gene Name | Fold Change | ye Direction | Pvalue |
|-----------------|------------|---|-------------|--------------|---------|
| rc_T16206_s_at | T16206 3 | 3065 EST | 4.29 | 음 | 0.00868 |
| rc_T16226_at | | 3066 EST | 7.23 | g | 0.00119 |
| rc_T16550_at | T16550 3 | 3072 vacuolar protein sorting 45B (yeast homolog) | 5.88 | d | 0.00004 |
| rc_T16652_s_at | T16652 3 | 3073 BCS1 (yeast homolog)-like | 3.63 | ਉ | 0.00434 |
| rc_T16983_s_at | T16983 3 | 3074 cleavage and polyadenylation specific factor 4, 30kD | 4.23 | đ | 0.0106 |
| rc_T17066_s_at | T17066 3 | 3075 SET domain, bifurcated, 1 | 5.14 | ဌ | 0.00073 |
| rc_T17339_f_at | T17339 3 | | 3.29 | đ | 0.00669 |
| rc_T17353_s_at | T17353 3 | | 3.52 | dn | 0.02085 |
| rc_T23426_s_at | T23426 3 | 3079 EST | 3.51 | g | 0.00674 |
| rc_T23465_at | T23465 3 | 3081 EST | 3.64 | d | 0.00265 |
| rc_T23516_f_at | T23516 3 | 3083 3-phosphoglycerate dehydrogenase | 3.39 | dn · | 0.00551 |
| rc_T24068_s_at | T24068 3 | 3088 EST | 8.65 | dn | 0.00118 |
| rc_T26471_at | T26471 3 | 3094 · EST | 3.76 | đ | 0.0165 |
| | | X-ray repair complementing defective repair in Chinese | Se | | |
| rc_T26646_f_at | T26646 3 | 3097 hamster cells 1 | 3.49 | dn | 0.02482 |
| rc_T30214_at | T30214 3 | 3099 EST | 4.46 | dn | 0.03654 |
| rc_T33489_s_at | T33489 3 | 3105 EST | 4.6 | dn | 0.00285 |
| rc_T33508_s_at | | _ | eta 4.96 | dn | 0.00064 |
| rc_T33619_s_at | | _ | 3.36 | dn | 0.01283 |
| rc_T33625_at | T33625 3 | 3108 EST | 3.36 | g | 0.04096 |
| rc_T33859_at | T33859 3 | 3109 KIAA0365 gene product | 3.9 | dn | 0.0019 |
| | | potassium voltage-gated channel, shaker-related | | | |
| T34377_at | T34377 3 | 3.111 subfamily, beta member 2 | 4.55 | đ | 0.00041 |
| rc_T40439_s_at | T40439 '3 | 3115 small nuclear ribonucleoprotein polypeptide B" | 3.06 | đ | 0.02842 |
| rc_T41078_at | T41078 3 | 3121 bromodomain adjacent to zinc finger domain, 2B | 3.08 | dn | 0.03426 |
| rc_T47032_s_at | T47032 3 | 3124 partner of RAC1 (arfaptin 2) | 60.9 | ဌ | 0.00019 |
| .rc_T47325_s_at | T47325 3 | 3125 EST | 5.6 | dn | 0.02923 |
| | | _ | | | - |
| rc_T47969_s_at | | | 3.19 | g | 0.02283 |
| rc_T51972_at | T51972 3 | 3140 EST | 3.44 | dn | 0.00406 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affv ID | Genbank Sed ID | ID Known Gene Name | Fold Change | Direction | Pvalue |
|-----------------------|----------------|--|-------------|-----------|---------|
| | | cytochrome P450, subfamily XIA (cholesterol side chain | | | |
| rc T53590 s at | T53590 3144 | 4 cleavage) | 5.09 | d | 0.00002 |
| rc T55196 at | T55196 3147 | | 4.24 | ф | 0.00046 |
| rc_T58607_at | | | 6.83 | dn | 0.03711 |
| rc T59668 s at | T59668 3160 | | 3.5 | dn | 0.00083 |
| rc T62521 at | | | 4.1 | dn | 0.00392 |
| rc_T62918_at | | _ | 5.25 | 롸 | 0.00687 |
| rc T65957 f at | T65957 3176 | '6 ribosomal protein S3A | 3.94 | dn | 0.04187 |
| rc T66935 at | | | 3.01 | dn | 0.00123 |
| rc T67053 f at | | 0 EST | ນ | dn | 0.01846 |
| rc_T77733 s at | | 19 tubulin, gamma 1 | 4 | ф | 0.00526 |
| rc_T78922_s_at | T78922 3222 | 22 stem cell growth factor; lymphocyte secreted C-type | 3.89 | ф | 0.00604 |
| rc_T79477 s at | T79477 3223 | | 4.4 | 롸 | 0.00074 |
| rc_T81393_s_at | | 28 HMT1 (hnRNP methyltransferase, S. cerevisiae)-like 1 | 3.32 | d | 0.0023 |
| rc T88814 at | | | 4.87 | d | 0.00001 |
| rc_T89703_at | | t4 EST | 5.27 | 롸 | 0.00019 |
| rc <u>T90190</u> s at | T90190 3247 | 47 H1 histone family, member 2 | 4.88 | 롸 | 0.00555 |
| rc T94452 at | | 57 EST | 3.23 | dn | 0.02245 |
| rc T95057 f at | | 59 EST | 6.46 | ф | 0.00613 |
| rc T97679 at | T97679 3266 | 36 EST | 3.32 | dn | 0.01566 |
| rc_T99312_at | T99312 3270 | 70 EST | 3.22 | dn | 0.00084 |
| U18018 at | U18018 3317 | 17 ets variant gene 4 (E1A enhancer-binding protein, E1AF) | 3.9 | ᅌ | 0.0403 |
| U18321_at | U18321 3318 | | 3.14 | 슠 | 0.00833 |
| ŀ | | proteasome (prosome, macropain) 26S subunit, non- | | | |
| U24704_at | U24704 3332 | | 3.45 | ф | 0.00037 |
| | | cyclin-dependent kinase inhibitor 2A (melanoma, p16, | | | |
| U26727 at | U26727 3337 | 37 inhibits CDK4) | 3.53 | d | 0.02913 |
| U30825_at | U30825 3343 | 43 splicing factor, arginine/serine-rich 9 | 3.07 | ᅀ | 0.01928 |
| U45285_at | U45285 3364 | 64 T-cell, immune regulator 1 | 5.75 | d | 0.00006 |
| U47025_s_at | U47025 3368 | 68 phosphorylase, glycogen; brain | 4.47 | ф | 0.00037 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank Seq ID | D Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|----------------|---|-------------|--------------|---------|
| | | siah binding protein 1; FBP interacting repressor; | | | |
| U51586_at | U51586 3386 | | 3.65 | 9 | 0.00403 |
| | | gamma-glutamyl hydrolase (conjugase, | | | |
| U55206_at | U55206 3392 | 2 folylpolygammaglutamyl hydrolase) | 3.34 | g | 0.00315 |
| U59321_at | U59321 3398 | 3 DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 17 | 3.12 | g | 0.02469 |
| U62392_at | U62392 3403 | •• | 3.56 | d | 0.0407 |
| U66661_at | U66661 3406 | - | 3.55 | d d | 0.0045 |
| U68142_at | U68142 3411 | | 3.02 | С | 0.0296 |
| | | SHC (Src homology 2 domain-containing) transforming | | | |
| U73377_at | U73377 3417 | _ | 4.64 | dn | 0.00081 |
| U73379_at | U73379 3418 | 8 ubiquitin carrier protein E2-C | 3.32 | d d | 0.00808 |
| U75285_rna1_at | U75285 3422 | 2 apoptosis inhibitor 4 (survivin) | 5.32 | 9 | 0.01127 |
| U85625_at | U85625 3448 | 8 ribonuclease 6 precursor | 4 | d | 0.01664 |
| U90426_at | U90426 3453 | 3 nuclear RNA helicase, DECD variant of DEAD box family | 3.24 | ဌ | 0.00035 |
| U90551_at | U90551 3457 | 7 H2A histone family, member L | 3.54 | 9 | 0.01523 |
| U90904_at | U90904 3458 | | 3.02 | g | 0.00381 |
| U91930_at | U91930 3460 | | 3.68 | d | 0.0000 |
| rc_W02041_at | W02041 3466 | | 5.34 | d | 0.00027 |
| rc_W02695_at | W02695 3467 | 7 EST | 3.1 | dn | 0.04745 |
| rc_W04507_s_at | W04507 3469 | 9 prefoldin 4 | 3.02 | d | 0.04091 |
| rc_W04550_at | W04550 3470 | EST | 4.01 | d | 0.00349 |
| rc_W15495_at | W15495 3474 | 4 chromosome 21 open reading frame 5 | 3.09 | d | 0.00491 |
| W26716_at | W26716 3482 | 2 non-histone chromosome protein 2 (S. cerevisiae)-like 1 | 5.47 | dn | 0.00146 |
| W28362_at | W28362 3488 | 8 KIAA0974 protein | 4.38 | dn | 0.00322 |
| W28366_at | W28366 3489 | 9 EST | 3.21 | d | 0.01007 |
| rc_W31906_at | W31906 3497 | 7 secretagogin | 6.62 | dn | 0.00926 |
| rc_W42627_f_at | W42627 3511 | | 4.37 | đ | 0.00021 |
| rc_W42674_at | W42674 3512 | 2 EST | 3.07 | đ | 0.0261 |
| rc_W42778_at | W42778 3513 | 3 EST | 3.27 | d d | 0.02411 |
| rc_W42788_at | W42788 351 | 4 deoxynucleotidyltransferase, terminal | 3.24 | 롸 | 0.02261 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank Sed ID | D Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|----------------|--|-------------|-----------|---------|
| rc W42957 at | W42957 3516 | s calmodulin 2 (phosphorylase kinase, delta) | 5.79 | d | 0.03669 |
| rc_W44557_at | W44557 3518 | 3 chromosome 1 open reading frame 2 | 4.1 | 슠 | 0.00433 |
| rc W45320 f at | W45320 3523 | 3 KRAB-associated protein 1 | 10.05 | dn | 0.00002 |
| rc W46286 s at | | _ | 3.68 | dn | 0.00311 |
| rc W46634 at | | EST | . 5.03 | 롸 | 0.02152 |
| rc W46810 s at | | 1 HMT1 (hnRNP methyltransferase, S. cerevisiae)-like 2 | 3.17 | d | 0.03434 |
| rc W46846 at | | EST | 5.11 | dn | 0.00025 |
| rc_W46947_at | | 3 EST | 3.42 | d | 0.04665 |
| rc_W47206_at | W47206 3535 | 5 EST | 3.23 | dn | 0.01931 |
| W49743 at | W49743 3540 | 0 EST | 3.11 | dn | 0.01121 |
| rc W49791 at | W49791 3541 | 1 plasminogen activator, tissue | 3.58 | dn | 0.02438 |
| rc_W56642_at | W56642 3547 | | 3.15 | ф | 0.00654 |
| rc_W57931_at | W57931 3549 | 9 EST | 3.01 | dn | 0.02661 |
| rc_W58081_at | W58081 3550 | 0 neuroendocrine-specific protein C like (foocen) | 3.05 | dn | 0.03767 |
| W58247 s at | W58247 3551 | 1 kinesin family member 4 | 3.08 | dn | 0.00048 |
| 1 | | DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide, Y | | | |
| rc W60097 at | W60097 3556 | 6 chromosome | 4.82 | dn | 0.04903 |
| rc_W60486_at | W60486 3558 | 8 EST | 4.27 | ф | 0.0046 |
| rc_W63608_at | W63608 3564 | 4 EST | 3.33 | ф | 0.02443 |
| W69302 at | W69302 3573 | 3 EST | 4.37 | g | 0.00165 |
| rc W69468 at | | 4 EST | 3.25 | g | 0.00055 |
| rc_W70336_at | W70336 3579 | _ | 4.46 | g, | 0.00023 |
| rc W72187 at | W72187 3582 | 2 EST | 3.09 | 핰 | 0.00134 |
| rc W72276 at | W72276 3583 | 3 EST | 3.56 | d | 0.0476 |
| rc_W73038_at | W73038 3588 | 8 EST | 4.83 | 롸 | 0.00902 |
| rc W74536 s at | W74536 3598 | 8 advanced glycosylation end product-specific receptor | 3.07 | 슠 | 0.00251 |
| rc_W78057_at | W78057 3600 | 0 EST | 5.53 | 롸 | 0.01231 |
| rc W79421 at | W79421 3603 | 3 EST | 3.57 | dn | 0.00895 |
| rc W79773 at | W79773 3605 | _ | 4.89 | 롸 | 0.00034 |
| rc_W80730_at | W80730 3607 | | 3.35 | dn | 0.01526 |
| rc_W80763_at | W80763 3608 | 8 EST | 4.98 | ф | 0.01026 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|------------------|-----------------|--------|--|-------------|----------------|---------|
| | | | KDEL (Lys-Asp-Glu-Leu) endoplasmic reticulum protein | | | |
| rc_W80852_at | W80852 | 3609 | retention receptor 3 | 6.37 | 9 | 0.00005 |
| . rc_W81654_at | W81654 | 3616 | SRY (sex determining region Y)-box 13 | 90.9 | đ | 0.00127 |
| rc_W84447_at | W84447 | 3617 | EST | 3.34 | g | 0.00986 |
| rc_W85875_at | W85875 | 3620 | EST | 4.91 | g | 0.01198 |
| rc_W86214_at | W86214 | 3625 | EST | 4.3 | g | 0.00194 |
| rc_W86748_at | W86748 | 3629 | EST | 5.09 | g | 0.01882 |
| rc_W90146_f_at | W90146 | 3644 | EST | 3.58 | <u> </u> | 0.00322 |
| rc_W92608_s_at | W92608 | 3653 | BAI1-associated protein 3 | 4.84 | g | 0.00599 |
| rc_W94281_s_at | W94281 | 3658 | integral membrane protein 2C | 3.51 | dn | 0.01689 |
| rc_W94885_at | W94885 | 3660 | EST | 6.53 | g | 0 |
| rc_W95841_at | W95841 | 3666 | EST | ဇ | ģ | 0.00466 |
| X04347_s_at | X04347 | 3680 | heterogeneous nuclear ribonucleoprotein A1 | 3.34 | d n | 0.00123 |
| X05610_at | X05610 | 3685 | collagen, type IV, alpha 2 | 6.04 | dn | 0.00062 |
| | | | collagen, type III, alpha 1 (Ehlers-Danlos syndrome type | | | |
| X06700_s_at | 00 2 90X | 3688 | IV, autosomal dominant) | 3.58 | dn | 0.02964 |
| | | | keratin 10 (epidermolytic hyperkeratosis; keratosis | | | |
| X14487_rna1_s_at | X14487 | 3702 | palmaris et plantaris) | 3.19 | dn | 0.01268 |
| X14850_at | X14850 | 3706 | H2A histone family, member X | 3.13 | <u>g</u> | 0.01523 |
| X17567_s_at | X17567 | 3719 | small nuclear ribonucleoprotein polypeptides B and B1 | 3.96 | g | 0.00317 |
| X53331_at | X53331 | 3727 | matrix Gla protein | 3.95 | d | 0.0151 |
| X54667_s_at | X54667 | 3731 | cystatin S, cystatin SN | 3.51 | dn | 0.00187 |
| X54941_at | X54941 | 3732 | CDC28 protein kinase 1 | 3.99 | d _n | 0.0016 |
| X54942_at | X54942 | 3733 | CDC28 protein kinase 2 | 3.8 | dn | 0.0035 |
| X56494_at | X56494 | 3738 | pyruvate kinase, muscle | 3.65 | d | 0.04795 |
| X57129_at | X57129 | 3743 | H1 histone family, member 2 | 4.63 | dn | 0.00663 |
| X57809_s_at | X57809 | 3746 | immunoglobulin lambda locus | 3.64 | dn | 0.02655 |
| X62153_s_at | X62153 | 3754 | minichromosome maintenance deficient (S. cerevisiae) 3 | | dn | 0.00704 |
| X62534_s_at | X62534 | 3755 | high-mobility group (nonhistone chromosomal) protein 2 | 3.39 | dn | 0.0186 |
| X64364_at | X64364 | 3764 | basigin | 3.57 | đ | 0.00902 |
| X66899_at | X66899 | 3772 | Ewing sarcoma breakpoint region 1 | 3.41 | dn | 0.03777 |
| | | | | | | |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change | Direction | Pvalue |
|----------------|---------|--------|---|-------------|--------------|---------|
| | | | transmembrane protein (63kD), endoplasmic | | | |
| X69910 at | X69910 | 3787 | reticulum/Golgi intermediate compartment | 3.35 | dn | 0.00898 |
| X74801 at | X74801 | 3791 | chaperonin containing TCP1, subunit 3 (gamma) | 3.86 | dn | 0.00453 |
| X79536 at | X79536 | 3804 | heterogeneous nuclear ribonucleoprotein A1 | 3.06 | dn | 0.00449 |
| X83425_at | X83425 | 3812 | Lutheran blood group (Auberger b antigen included) | 3.66 | d | 0.02661 |
| X87212 at | X87212 | 3816 | cathepsin C | 3.45 | g | 0.02486 |
| I | | | DNA segment on chromosome X (unique) 9879 | | | |
| X92896 at | X92896 | 3829 | expressed sequence | 3.1 | 롸 | 0.0405 |
| X99920_at | X99920 | 3843 | S100 calcium-binding protein A13 | 4.66 | d | 0.00113 |
| Y00705 at | Y00705 | 3850 | serine protease inhibitor, Kazal type 1 | 28.88 | 롸 | 0.00003 |
| Y00764_at | Y00764 | 3851 | ubiquinol-cytochrome c reductase hinge protein | 3.04 | dn | 0.01294 |
| Y08302_at | Y08302 | 3852 | dual specificity phosphatase 9 | 3.48 | dn | 0.00787 |
| Y08999_at | Y08999 | 3855 | actin related protein 2/3 complex, subunit 1A (41 kD) | 3.13 | dn | 0.02376 |
| I | | | eukaryotic translation elongation factor 1 delta (guanine | | | |
| Z21507 at | Z21507 | 3867 | nucleotide exchange protein) | 3.57 | dn | 0.01898 |
| Z23090_at | ZZ3090 | 3868 | heat shock 27kD protein 1 | 3.69 | dn | 0.00628 |
| Z24727 at | Z24727 | 3871 | tropomyosin 1 (alpha) | 3.64 | dn | 0.00388 |
| Z37987 s at | Z37987 | 3882 | glypican 3 | 10.66 | dn | 0.02304 |
| rc Z38299 at | Z38299 | 3887 | EST | 3.71 | dn | 0.0036 |
| rc_Z38431_at | Z38431 | 3889 | EST | 3.09 | dn | 0.0083 |
| rc_Z38444_at | Z38444 | 3891 | KIAA0923 protein | 3.38 | dn | 0.02918 |
| rc Z38462 at | Z38462 | 3892 | KIAA0938 protein | 4.69 | dn | 0.0142 |
| rc_Z38904_at | Z38904 | 3896 | EST | 3.02 | dn | 0.00814 |
| rc_Z39191_at | Z39191 | 3901 | EST | 3.47 | dn | 0.00756 |
| rc_Z39200_at | Z39200 | 3902 | EST | 4.07 | đ | 0.00075 |
| rc_Z39379_at | Z39379 | 3903 | EST | 3.77 | g. | 0.00513 |
| rc_Z39429_at | Z39429 | 3906 | EST | 3.58 | dn | 0.00416 |
| rc Z39682 s at | Z3968Z | 3911 | KIAA0954 protein | 96.9 | dn | 0.01966 |
| rc_Z40006_at | Z40006 | 3917 | EST | 3.54 | dn | 0.00156 |
| rc_Z40883_r_at | Z40883 | 3924 | EST | 3.26 | 앜 | 0.01863 |
| rc Z41349 at | Z41349 | 3931 | EST | 3.45 | d | 0.01503 |

Table 7B. Down regulated in hepatocellular carcinoma versus normal sample set 2.

| Affy ID | Genbank | Seq ID | Known Gene Name | Fold Change Direction Pvalue | Direction | Pvalue |
|--------------|---------|--------|--|------------------------------|-----------|---------|
| | | | succinate dehydrogenase complex, subunit A, | | | |
| rc_Z41747_at | Z41747 | 3936 | flavoprotein (Fp) | 3.04 | g | 0.01336 |
| Z47727_at | Z47727 | 3940 | polymerase (RNA) II (DNA directed) polypeptide K | 3.28 | d | 0.00317 |
| Z74615_at | Z74615 | 3949 | collagen, type I, alpha 1 | 5.47 | d | 0.00283 |
| Z74616_s_at | Z74616 | 3950 | collagen, type I, alpha 2 | 5.95 | dn | 0.02212 |

Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| | | | | fold | | hcc | hcc | hcc | | | |
|----------------|----------|--------|---|--------|--------------|--------|--------|---------|----------------|--------|---------|
| | | | | change | | sample | samble | sample | normal | normal | normal |
| | | | | in hcc | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | set 2 | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | DEK oncogene (DNA | | | | | | | | |
| AA093497 s at | AA093497 | 199 | binding) | #N/A | #N/A | 180.58 | 107.64 | 165.71 | 52.11 | 48.84 | 26.51 |
| AA248283 at | AA248283 | 534 | EST | #N/A | #N/A | 67.18 | 45.86 | 55.18 | 21.63 | 18.96 | 11.71 |
| AA291456 s at | AA291456 | 200 | EST | #N/A | #N/A | 828.34 | 830.24 | 196.63 | 671.21 | 509.58 | 629.13 |
| | | | GAS2-related on | | | | | | | | |
| AA400643 s at | AA400643 | 817 | chromosome 22 | 4.04 | 0.03751 | 118.72 | 123.18 | 137.15 | -43.6 | -33.96 | 60.39 |
| AA421079 at | AA421079 | 930 | EST | #N/A | #N/A | 37.17 | 34.26 | 18.5 | 15.27 | 16.75 | 15.28 |
| ı | | | Notch (Drosophila) homolog | | | | | | | | |
| -AA428172 f at | AA428172 | 986 | က | 9.63 | 0.00195 | 335.57 | 374.9 | 231.52 | -9.64 | -15.61 | 56.16 |
| AA464043 s at | AA464043 | 1255 | EST | 3.99 | 0.00056 | 116.97 | 115.99 | 53.49 | 14.27 | 10.06 | 25.86 |
| H19562 at | H19562 | 1896 | PTD010 protein | #N/A | #N/A | 522.13 | 432.65 | 256.94 | 393.61 | 326.53 | 182.71 |
| L08044 s at | L08044 | 2149 | trefoil factor 3 (intestinal) | #N/A | #N/A | 214.6 | 108.17 | 238.79 | 107.82 | 58.55 | 184.16 |
| L29218 s at | L29218 | 2190 | CDC-like kinase 2 | 6.51 | 0.00019 | 219.64 | 237.35 | 104.29 | 6.53 | -10.03 | 59.8 |
| L37747 s at | L37747 | 2206 | lamin B1 | #N/A | #N/A | 43.42 | 28.65 | 37.33 | 5.97 | 5.1 | 7.57 |
| N34257 at | N34257 | 2495 | EST | #N/A | #N/A | 20.76 | 21.48 | 19.54 | -1.84 | -2.74 | 8.53 |
| R50692 at | R50692 | 2888 | KIAA0476 gene product | #N/A | #N/A | 197.89 | 187.87 | 101.08 | 148.53 | 151.12 | 63.03 |
| R60368_at | R60368 | 2917 | EST | #N/A | #N/A | 29.68 | 28.93 | 31.9 | -28.29 | -26.44 | 41.68 |
| R66475 at | R66475 | 2938 | EST | #N/A | #N/A | 45.68 | 32.34 | 34.37 | 7.64 | 8.26 | 6.3 |
| l | | | potassium voltage-gated channel, shaker-related | | | | | | | | |
| T34377_at | T34377 | 3111 | subfamily, beta member 2 DEAD/H (Asp-Glu-Ala- | 4.55 | 0.00041 | 113.59 | 120.58 | 51.65 | 6.87 | 11.17 | 25.69 |
| | | | Asp/His) box polypeptide 11 (S.cerevisiae CHL1-like | | | | | | | | |
| U75968_s_at | U75968 | 3423 | helicase) | ¥/\# | #N/A #N/A | 187.33 | 200.16 | 90.35 | 115.5 45.59 | 102.24 | 77.72 |
| W28696_i_at | W28696 | 3491 | EST | #N/A | W/W# | 14.7 | | 17.12 | | | 2.47 |

Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| | | | | told change | | hcc sample | hcc sample | hcc sample | normal | normai | normal |
|------------------|----------|--------|--------------------------------|----------------|---------|---------------|---------------|---------------|--------|--------|---------|
| | | | | in hcc | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | set 2 | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | H1 histone family, member | | | | | | | | |
| X57129_at | X57129 | 3743 | 2 | 4.63 | 0.00663 | 137.13 | 137.75 | 87.31 | 8.38 | 5.39 | 28.3 |
| rc AA024658 at | AA024658 | 47 | ribosomal protein S19 | 7.55 | 0.00592 | 278.11 | 120.83 | 373.3 | 7.62 | 5.03 | 10.68 |
| rc AA029288 at | AA029288 | 65 | EST | 3.36 | 0.04908 | 134.53 | 56.15 | 209.69 | 4.09 | 3.35 | 8.39 |
| rc_AA037828_at | AA037828 | 88 | KIAA0614 protein | #N/A | #N/A | 53.61 | 46.86 | 41.68 | 13.32 | 10.65 | 11.96 |
| rc_AA040465_at | AA040465 | 92 | EST | 3.63 | 0.01806 | 229.8 | 210.67 | 180.94 | 47.15 | 45.53 | 5.48 |
| rc_AA053660_at | AA053660 | 128 | EST | #N/A | #N/A | 543.66 | 223.47 | 685.44 | 95.24 | 79.97 | 42.99 |
|) | | | cAMP responsive element | | | | | | | | |
| rc_AA136332_at | AA136332 | 299 | binding protein 3 (luman) | #N/A | #N/A | 49.04 | 33.07 | 31.07 | -18.27 | -20.23 | 13.05 |
| rc_AA149530_at | AA149530 | 324 | interferon regulatory factor 3 | #N/A | #N/A | 32.94 | 23.12 | 34.23 | 4.21 | 5.38 | 8.53 |
| rc_AA149586_at | AA149586 | 325 | EST | #N/A | #N/A | 24.99 | 26.6 | 34.37 | 5.26 | 9.11 | 23.41 |
| rc_AA206023_at | AA206023 | 427 | EST | #N/A | #N/A | 154.13 | 170.36 | 85.33 | 108.57 | 111.76 | 58.39 |
| ! | | | N-ethylmaleimide-sensitive | | | ٠ | | | | | |
| rc_aa234530_s_at | AA234530 | 484 | factor | #N/A | #N/A | 143.67 | 135.48 | 82.59 | 51.33 | 42.9 | 35.02 |
| rc_AA251909_at | AA251909 | 549 | EST | 3.59 | 0.01129 | 92.19 | 79.34 | 72.73 | 8.88 | 11.22 | 8.27 |
| rc_AA262030_at | AA262030 | 605 | EST | #N/A | #N/A | 119.75 | 89.66 | 103.84 | 31.51 | 32.91 | 14.46 |
| i | - | ` | ribonuclease HI, large | | | | | | | | |
| rc_AA262477_at | AA262477 | 809 | subunit | 4.57 | 0.00724 | 242.42 | 199.5 | 182.26 | 44.51 | 36.87 | 22.37 |
| rc_AA283759_at | AA283759 | 671 | EST | #N/A | #N/A | 91.82 | 76.73 | 42.28 | 46.87 | .45.86 | 21.54 |
| rc_AA291644_at | AA291644 | 701 | EST | 3.28 | 0.00033 | 114.73 | 117.29 | 44.59 | 32.77 | 30.24 | 21.1 |
| rc_AA338760 at | AA338760 | 744 | EST | 3.96 | 0.01307 | 129.77 | 130.26 | 87.27 | 14.91 | 21.91 | 26.96 |
| rc_AA398205_at | AA398205 | 789 | EST | 4.22 | 0.00059 | 125.4 | 107.84 | 63.85 | 15.01 | 7.37 | 26.26 |
| | | | tumor suppressor deleted in | | | | | | | | |
| rc_AA401965_at | AA401965 | 833 | oral cancer-related 1 | 7.58 | 0.00089 | 932.74 | 924.02 | 593.69 | 120.58 | 93.29 | 109.41 |
| rc_AA402272_at | AA402272 | 837 | EST | 3.73 | 0.02336 | 348.91 | 340 | 230.15 | 99.7 | 92.28 | 80.27 |
| rc_AA404597_at | AA404597 | 854 | EST | #N/A | #N/A | 609.98 | 525.02 | 371.31 | 379.26 | 336.33 | 167.43 |
| rc_AA417030_at | AA417030 | 914 | EST | 7.35 | 0.00555 | 200.3 | 168.21 | 135.63 | 6.42 | 7.5 | 7.77 |

Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| | | | | fold | | pcc | hcc | hcc | | | |
|------------------|----------|--------|-------------------------------|--------|---------|--------|--------|---------|--------|--------|---------|
| | | | | 0000 | | olumes | olumes | eamnle | normal | normal | normai |
| | | | | cnange | | samble | sample | Sample | 10 I | | |
| | | | | in hcc | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| Affv ID | Genbank | Sea ID | Known Gene Name | set 2 | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| re AA423820 at | AA423820 | 940 | EST | #N/A | #N/A | 111.38 | 111.23 | 120.84 | 41.9 | 21.93 | 44.58 |
| rc AA425852 i at | AA425852 | 928 | EST | 3.82 | 0.0395 | 131.52 | 95.35 | 166.76 | 7.02 | 5.22 | 4.75 |
| 1 | | | pituitary tumor-transforming | | | | | | | | |
| rc AA430032 at | AA430032 | 1009 | · | 10.67 | 0.00052 | 377.69 | 320.44 | 294.49 | 26.8 | 15.31 | 35.24 |
| rc_AA430673_at | AA430673 | 1017 | EST | #N/A | #N/A | 45.74 | 34.15 | 44.8 | -1.21 | 3.63 | 15.26 |
| rc_AA443321_at | AA443321 | 1076 | EST | #N/A | #N/A | 33.93 | 32.89 | 11.76 | 10.79 | 9.65 | 7.26 |
| | | | tumor suppressing | | | | | | | | |
| rc AA443941 at | AA443941 | 1085 | subtransferable candidate 1 | #N/A | W/V# | 97.74 | 103.38 | 47.48 | 31.09 | 34.9 | 16.61 |
| rc_AA447223_at | AA447223 | 1100 | EST | #N/A | #N/A | 31.66 | 18.91 | 31.1 | 4.45 | 3.74 | 6.5 |
| 1 | | ٠ | translation initiation factor | | | | | | | | |
| rc aa449431 s at | AA449431 | 1124 | IF2 | 3.76 | 0.00571 | 94.41 | 95.64 | 54.08 | 17.4 | | 5.38 |
| rc AA454597 s at | AA454597 | 1166 | EST | 4.23 | 0.00917 | 342.59 | 311.81 | 207.71 | 9.99 | | 23.41 |
| rc AA45887 at | AA45882 | 1205 | EST | #N/A | #N/A | 93.56 | 52.28 | 98.72 | 24.97 | 22.52 | 17.66 |
| rc_AA458934_at | AA458934 | 1208 | EST | #N/A | #N/A | 33.6 | 28.19 | 29.02 | 7.1 | 7.91 | 4.22 |
| rc AA465347 at | AA465342 | 1271 | EST | 3.21 | 0.01378 | 131.07 | 109.99 | 95.4 | 30.07 | 33.06 | 23.97 |
| rc_AA477561_at | AA477561 | 1292 | EST | #N/A | #N/A | 146.3 | 127.03 | 97.99 | 63.83 | 76.56 | 37.87 |
| 1 | | | calcium/calmodulin- | | | | | | | | |
| | | | dependent protein kinase | | | | | | | | 1 |
| rc AA491295 at | AA491295 | 1390 | kinase 2, beta | 3.71 | 0.0103 | 180.61 | 173.47 | 118.85 | 46.95 | 31.51 | 65.56 |
| Ī | | | RAD51 (S. cerevisiae) | | | | | | | | |
| | | | homolog (E coli RecA | | | | | | | | |
| rc AA608545 at | AA608545 | 1462 | homolog) | W/A | #N/A | 70.5 | 57.51 | 53.36 | 7.76 | 8.92 | 6.17 |
| 1 | | | flap structure-specific | | | | | | | | |
| rc AA620553 s at | AA620553 | 1504 | endonuclease 1 | 7.56 | 0.00101 | 262.87 | 232.63 | (1) | | | |
| rc AA621325 at | AA621325 | 1522 | HNK-1 sulfotransferase | #N/A | #N/A | 74.57 | 70.64 | | | | • |
| rc_AA621780_at | AA621780 | 1530 | CGI-96 protein | #N/A | #N/A | 81.72 | 92.23 | | | | |
| rc F02807 at | F02807 | 1781 | KIAA0838 protein | 5.67 | 0.02064 | 260.02 | 219.01 | 229.96 | 20.93 | 16.66 | 22.24 |
| | ; | | - | | | | | | | | |

Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| | | | | fold | | hcc | hcc | pcc | | | |
|----------------|---------|--------|--------------------------|--------|---------|--------|--------|---------|--------|--------|---------|
| | | | | change | | sample | sample | samble | normal | normal | normal |
| | | | | in hcc | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | set 2 | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| rc_F04444_at | F04444 | 1788 | EST | 4.13 | 0.00944 | 371.21 | 360.5 | 186.45 | 119.68 | 121.27 | 109.82 |
| rc_F04524_s_at | F04524 | 1790 | stomatin-like protein 1 | W/A | #N/A | 117.77 | 135.65 | 68.29 | 46.56 | 44.53 | 33.75 |
| rc_F08876_at | F08876 | 1797 | EST | 9.06 | 0 | 209.64 | 201.85 | 53.21 | -22.5 | -20.17 | 37.21 |
| rc_F10161_at | F10161 | 1811 | EST | #N/A | #N/A | 50.78 | 46.73 | 32.42 | 15.4 | 14.78 | 11.87 |
| rc_F10453_at | F10453 | 1819 | EST | 3.64 | 0.01878 | 135.07 | 138.53 | 85.7 | 27.28 | 14.2 | 54.04 |
| rc_H04649_at | H04649 | 1838 | EST | #N/A | #N/A | 180.25 | 158.18 | 164.34 | 50.86 | 41.2 | 33.9 |
| rc_H04793_at | H04793 | 1840 | DKFZP434F091 protein | #N/A | #N/A | 23.18 | 20 | 41.25 | -14.45 | -20.45 | 29 |
| rc_H59617_at | H59617 | 1964 | EST | 5.81 | 0.0115 | 212.68 | 209.19 | 157.42 | 19.24 | 21.65 | 16.88 |
| rc_H97012_at | H97012 | 2058 | EST | 3.51 | 0.03505 | 171.34 | 138.12 | 170.68 | 27.69 | 38.46 | 36.08 |
| rc_H98657_at | H98657 | 2068 | EST | #N/A | #N/A | 141.17 | 131.42 | 69.63 | 111.2 | 84.74 | 128.92 |
| rc_N23319_at | N23319 | 2454 | EST | #N/A | #N/A | 70.62 | 66.39 | 49.4 | 13.74 | 8.56 | 21.62 |
| rc_N23868_at | N23868 | 2459 | EST | #N/A | #N/A | 79.69 | 81.67 | 56.06 | 20.52 | 20.19 | 11.48 |
| rc_N45224_at | N45224 | 2516 | EST | #N/A | #N/A | 349.73 | 360.67 | 164.87 | 215.49 | 223.16 | 151.06 |
| rc_N48595_at | N48595 | 2528 | EST | #N/A | #N/A | 85.77 | 67.62 | 51.78 | 28.84 | 29.38 | 8.3 |
| rc_N75541_at | N75541 | 2692 | EST | 4.43 | 0.01059 | 182.83 | 131.35 | 183.41 | 34.1 | 15.82 | 56.07 |
| | | | ras homolog gene family, | | | | | | | | |
| rc_N90273_s_at | N90273 | 2713 | member H | #N/A | #N/A | 105.53 | 96.01 | 41.89 | | 59.56 | 35.69 |
| rc_n93465_s_at | N93465 | 2735 | EST | #N/A | W/V# | 803.68 | | 385.5 | 624.57 | 694.14 | 306.11 |
| ۰′۱ | R39191 | 2834 | KIAA1020 protein | 5.18 | 0.03185 | 331.72 | | 590.96 | | 18.33 | 16.04 |
| rc_R44793_at | R44793 | 2859 | EST | 5.4 | 0.00329 | 155.86 | | 125.26 | | 7.86 | 16.54 |
| rc_R44817_at | R44817 | | EST . | #N/A | W/V# | 594.46 | | 206.15 | | 249.29 | 60.69 |
| rc_R44896_at | R44896 | 2862 | KIAA0665 gene product | #N/A | #N/A | 67.08 | | 34.18 | | 20.75 | 23.04 |
| rc_R64137_s_at | R64137 | 2931 | EST | #N/A | W/N# | 31.25 | | 21 | | 2.65 | 6.94 |
| rc_R71395_at | R71395 | 2952 | EST | 4.12 | 0.03719 | 166.03 | | 190.98 | | 12.28 | 14.58 |
| rc_R72087_at | R72087 | 2955 | EST | #N/A | #N/A | 262.11 | | 92 | | 183.76 | 83.41 |
| rc_R91819_at | R91819 | 2984 | EST | 12.81 | 0.00037 | 467.79 | | 426.16 | | 9.35 | 36.49 |
| rc_T03749_at | T03749 | 3048 | KIAA1089 protein | 4.23 | 0.00776 | 103 | | 74.34 | | -6.08 | 13.69 |
| rc_T25744_s_at | T25744 | 3092 | EST | #N/A | #N/A | 72.74 | | 37.84 | | 12.65 | 13.18 |

· Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| Cenbank Seq ID Known Gene Name change in hcc set 2 in hcc set 2 in complexed in hcc set 2 inc metalloproteinase, and set 3169 EST F.25 inc metalloproteinase, inches set 2 inc metalloproteinase, and set 316 set 2 inc metalloproteinase, and set 316 set 321 set 321 set 321 set 322 set 322 set 322 set 322 set 322 set 322 set 332 set 333 set 3333 set 3333 set 333 set 333 set 3333 set 333 set 333 set 3333 set 333 set 333 set 333 set 333 | | fold | | hcc | hcc | pcc | | | |
|--|-----------------------|--------------|------------|--------|--------|---------|--------|--------|----------|
| In hcc Genbank Seq ID Known Gene Name Set 2 F251 T62918 3169 EST EST T62918 3169 EST T62918 3169 EST T62918 3169 EST T62918 3169 EST Tinc metalloproteinase, and rabb GTPase activating Trabb GTPase activation Trabb GTPase activating Trabb GTPase activating Trabb GTPase activating Trabb GTPase activating Trabb GTPase activation Trabb GTPase activating Trabb GTPase activation Trabb GTPase activat | | chanc | je Je | samble | sample | sample | normal | normal | normal |
| Genbank Seq ID Known Gene Name set 2 T62918 3169 EST 5.25 T62918 3169 EST #N/A T89731 3245 EST #N/A zinc metalloproteinase, #N/A #N/A rab6 GTPase activating #N/A protein (GAP and #N/A w47388 3536 centrosome-associated) #N/A w78075 3608 EST 4.98 w80763 3608 EST 4.98 w80763 3608 EST 4.98 w80764 EST 4.98 w80763 3892 EST 4.98 x38462 3893 EST #W/A x38688 3893 EST #W/A x38699 3894 EST #W/A x38729 3894 EST #W/A x40556 3921 CGI-96 protein #W/A x4 AA02659 135 collagen, type V, alpha 1 10.87 | | in hc | ຸ ບ | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| T62918 | | | 2 p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| T89731 3245 EST #N/A zinc metalloproteinase, and waz176 3498 STE24 (yeast, homolog) #N/A rab6 GTPase activating protein (GAP and w47388 3536 centrosome-associated) #N/A w78057 3600 EST 5.53 w80763 3608 EST 7.38299 3887 EST 7.38404 3888 EST 7.38404 3888 EST 7.38404 3898 EST 7.38404 3898 EST 7.38405 3921 CGI-96 protein #N/A z40556 3921 CGI-96 protein #N/A z405696 135 collagen, type V, alpha 1 10.87 at AA012979 225 vaccinia related kinase 1 #N/A at AA056914 428 EST #N/A #N/A at AA206914 428 EST #N/A #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243598 531 EST #N/A #N/A at AA243598 531 EST #N/A #N/A #N/A #N/A #N/A #N/A #N/A #N/A | | 5. | 25 0.00687 | 136.22 | | 83.63 | -12.41 | -6.89 | 17.59 |
| zinc metalloproteinase, #N/A rab6 GTPase activating protein (GAP and W47388 3536 centrosome-associated) #N/A W80763 3608 EST 4.98 W80763 3608 EST 4.98 W80763 3608 EST 4.98 W80763 3608 EST 4.98 Z38299 3887 EST 3.71 Z38462 3892 KIAA0938 protein 4.69 Z38729 3894 EST #N/A Z38729 3894 EST #N/A Z38729 3894 EST #N/A Z38729 3894 EST #N/A Z40566 3921 CGI-96 protein #N/A Z4056 3924 EST #N/A AA021549 42 EST #N/A at AA112979 225 vaccinia related kinase 1 #N/A at AA134063 286 EST #N/A at | | #N/# | W/W# | 63.59 | 65.07 | 20.47 | 5.75 | 3.83 | 11.8 |
| #N/A rab6 GTPase activating protein (GAP and wa7388 3536 centrosome-associated) #N/A w77388 3536 centrosome-associated) #N/A w77383 3536 centrosome-associated) #N/A w80763 3608 EST 4.98 w80763 3608 EST 3.73 238404 3888 EST #N/A z38462 3892 KIAA0938 protein 4.69 z38688 3893 EST #N/A z40556 3921 CGI-96 protein #N/A z40556 3921 CGI-96 protein #N/A z4056 3921 CGI-96 protein #N/A z40569914 428 EST #N/A #N/A at AA243133 525 serine/threonine kinase 15 #N/A at AA243133 525 serine/threonine kinase 15 #N/A at AA243598 531 EST #N/A #N/A #N/A at AA243598 531 EST #N/A #N/A #N/A #N/A #N/A #N/A #N/A #N/A | zinc metalloproteinas | | | | | | | | ! |
| rab6 GTPase activating protein (GAP and W47388 3536 centrosome-associated) #N/A W78057 3600 EST 4.98 W80763 3608 EST 3.58 EST 238299 3887 EST 3.71 Z38404 3888 EST #N/A Z38462 3892 KIAA0938 protein 4.69 Z3868 3893 EST #N/A Z38729 3894 EST #N/A Z40556 3921 CGI-96 protein #N/A Z40569 135 collagen, type V, alpha 1 10.87 at AA026270 56 tyrosyl-tRNA synthetase #N/A at AA056896 135 collagen, type V, alpha 1 10.87 at AA12979 225 vaccinia related kinase 1 #N/A at AA206914 428 EST #N/A #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA24366 527 EST #N/A #N/A at AA243598 531 EST #N/A #N/A at AA243598 531 EST #N/A #N/A #N/A #N/A #N/A #N/A #N/A #N/A | - | | W/A | 550.53 | 614.92 | 234.79 | 290.18 | 277.95 | 129.45 |
| W47388 3536 centrosome-associated) #N/A W78057 3600 EST 5.53 W80763 3608 EST 4.98 W80746 3644 EST 3.58 Z38299 3887 EST 3.71 Z38462 3892 KIAA0938 protein 4.69 Z38688 3893 EST #N/A Z38729 3894 EST #N/A Z4056 3921 CGI-96 protein #N/A Z4056 3921 CGI-96 protein #N/A AA021549 42 EST #N/A at AA026570 56 tyrosyl-tRNA synthetase #N/A at AA112979 225 vaccinia related kinase 1 #N/A at AA134063 286 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243598 531 EST #N/A at AA243598 531 EST #N/A< | rab6 GTPase activatii | Ð. | | | | | | | |
| W78057 3600 EST 5.53 W80763 3608 EST 4.98 W80746 3644 EST 3.58 Z38299 3887 EST 4.98 Z38462 3892 KIAA0938 protein 4.69 Z3868 3893 EST #N/A Z38729 3894 EST #N/A Z4056 3921 CGI-96 protein #N/A AA021549 42 EST #N/A AA026270 56 tyrosyl-tRNA synthetase #N/A at AA056896 135 collagen, type V, alpha 1 10.87 at AA12979 225 vaccinia related kinase 1 #N/A at AA15063 286 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243466 527 EST #N/A at AA243598 531 | | _ | Y/N# | 97.33 | 71 | 59.73 | 43.93 | 36.52 | 48.34 |
| W80763 3608 EST 4.98 W807163 3608 EST 3.58 Z38299 3887 EST 3.71 Z38462 3892 KIAA0938 protein 4.69 Z3868 3893 EST #N/A Z38729 3894 EST #N/A Z40556 3921 CGI-96 protein #N/A AA021549 42 EST #N/A AA026270 56 tyrosyl-tRNA synthetase #N/A at AA056896 135 collagen, type V, alpha 1 10.87 at AA12979 225 vaccinia related kinase 1 #N/A at AA134063 286 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243466 527 EST #N/A at AA243598 531 EST #N/A at AA243598 | | | 3 | 265.23 | 222.35 | 199.24 | 29.21 | 29.9 | 34.33 |
| tf W90146 3644 EST 3.58 Z38299 3887 EST 3.71 Z38404 3888 EST #N/A Z38462 3892 KIAA0938 protein 4.69 Z38729 3894 EST #N/A Z40556 3921 CGI-96 protein #N/A at AA021549 42 EST #N/A at AA026270 56 tyrosyl-tRNA synthetase #N/A at AA056896 135 collagen, type V, alpha 1 10.87 at AA112979 225 vaccinia related kinase 1 #N/A at AA134063 286 EST #N/A at AA206914 428 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243466 527 EST #N/A at AA243598 531 EST | | 4 | _ | 147.37 | 138.37 | 122.38 | 16.84 | 15.48 | |
| 238299 3887 EST 3.71 238404 3888 EST #N/A 238462 3892 KIAA0938 protein 4.69 238729 3894 EST #N/A 240556 3921 CGI-96 protein #N/A 240556 3921 CGI-96 protein #N/A 317 AA026270 56 tyrosyl-tRNA synthetase #N/A 317 AA056896 135 collagen, type V, alpha 1 10.87 317 AA12979 225 vaccinia related kinase 1 #N/A 327 EST #N/A #N/A 327 EST #N/A 327 EST #N/A 328 EST #N/A 327 EST #N/A 328 EST #N/A 327 EST #N/A 328 EST #N/A 329 Serine/threonine kinase 15 7.03 327 EST #N/A 328 | | က် | | 82.11 | 69.02 | 47.2 | 9.93 | 8.63 | |
| 238404 3888 EST #N/A 238462 3892 KIAA0938 protein 4.69 238688 3893 EST #N/A 238729 3894 EST #N/A 240556 3921 CGI-96 protein #N/A at AA021549 42 EST 3.17 at AA026270 56 tyrosyl-tRNA synthetase #N/A at AA012979 225 vaccinia related kinase #N/A at AA112979 225 vaccinia related kinase #N/A at AA14063 286 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 7.03 at AA243466 527 EST #N/A at AA243598 531 EST #N/A at AA243598 531 EST #N/A | | က် | | _ | 127.16 | | 30.48 | 25.94 | |
| Z38462 3892 KIAA0938 protein 4.69 Z38688 3893 EST #N/A Z40556 3921 CGI-96 protein #N/A AA021549 42 EST #N/A at AA026570 56 tyrosyl-tRNA synthetase #N/A at AA112979 225 vaccinia related kinase 1 #N/A at AA112979 225 vaccinia related kinase 1 #N/A at AA134063 286 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243358 531 EST #N/A at AA243588 531 EST #N/A at AA243588 531 EST #N/A | | //N# | | | 56.71 | | 29.82 | 30.16 | |
| Z38688 3893 EST #N/A Z40556 3924 EST #N/A Z40556 3921 CGI-96 protein #N/A At A0021549 42 EST 3.17 At A0026270 56 tyrosyl-tRNA synthetase #N/A At A0056896 135 collagen, type V, alpha 1 10.87 At A412979 225 vaccinia related kinase 1 #N/A At A4134063 286 EST #N/A At A4150053 327 EST #N/A AA206914 428 EST #N/A AA243133 525 serine/threonine kinase 15 7.03 AA243358 531 EST #N/A AA243598 531 EST #N/A AA252147 551 EST #N/A | _ | 4 | | | | 116.16 | -0.86 | 0.53 | |
| Z38729 3894 EST #N/A Z40566 3921 CGI-96 protein #N/A at AA021549 42 EST 3.17 at AA026270 56 tyrosyl-tRNA synthetase #N/A at AA112979 225 vaccinia related kinase 1 #N/A at AA134063 286 EST #N/A at AA150053 327 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243166 527 EST #N/A at AA243598 531 EST #N/A at AA243598 531 EST #N/A | | /N# | | | | | 21.38 | 13.4 | |
| Z40556 3921 CGI-96 protein #N/A at AA021549 42 EST 3.17 at AA026270 56 tyrosyl-tRNA synthetase #N/A at AA055896 135 collagen, type V, alpha 1 10.87 at AA112979 225 vaccinia related kinase 1 #N/A at AA134063 286 EST #N/A at AA150053 327 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243566 527 EST #N/A at AA243598 531 EST #N/A at AA243598 531 EST #N/A | | /N# | | 77.67 | 57.81 | | -2.4 | -5.01 | 46.96 |
| at AA021549 42 EST 3.17 at AA026270 56 tyrosyl-tRNA synthetase #N/A at AA055896 135 collagen, type V, alpha 1 10.87 at AA112979 225 vaccinia related kinase 1 #N/A at AA134063 286 EST #N/A at AA150053 327 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243566 527 EST #N/A at AA243598 531 EST #N/A at AA243598 531 EST #N/A | | /N# | | | | | | | |
| at AA055896 135 collagen, type V, alpha 1 10.87 at AA12979 225 vaccinia related kinase 1 #N/A at AA134063 226 vaccinia related kinase 1 #N/A at AA134063 327 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243598 531 EST #N/A #N/A at AA243598 531 EST #N/A #N/A at AA243598 531 EST #N/A #N/A #N/A #N/A #N/A #N/A #N/A #N/A | | ю́ | | | | | | | |
| at AA055896 135 collagen, type V, alpha 1 10.87 at AA112979 225 vaccinia related kinase 1 #N/A at AA134063 226 EST #N/A #N/A at AA206914 428 EST #N/A #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA24366 527 EST #N/A at AA243598 531 EST #N/A #N/A at AA243598 531 EST #N/A #N/A #N/A #N/A #N/A #N/A #N/A #N/A | | | | | | 191.51 | 76.85 | | |
| at AA112979 225 vaccinia related kinase 1 #N/A at AA134063 286 EST #N/A at AA150053 327 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243598 531 EST #N/A at AA243598 531 EST #N/A at AA252147 551 EST #N/A #N/A | _ | - | | | | | | | |
| at AA134063 286 EST #N/A at AA150053 327 EST #N/A at AA206914 428 EST #N/A at AA243133 525 serine/threonine kinase 15 7.03 at AA243566 527 EST #N/A at AA243598 531 EST #N/A at AA252147 551 EST #N/A | | _ | | | | | | | |
| at AA243143 527 EST #N/A AA243143 525 serine/threonine kinase 15 T.03 at AA243598 531 EST #N/A AA243598 531 EST #N/A AA252147 551 EST #N/A #N/A | | | | | | | | | |
| at AA24314 428 EST #N/A at AA24313 525 serine/threonine kinase 15 7.03 at AA243598 531 EST #N/A at AA252147 551 EST #N/A #N/A | | N# | | | | • | • | | 73.73 |
| at AA243143 525 serine/threonine kinase 15 7.03 at AA243598 531 EST #N/A at AA252147 551 EST #N/A | | N# | | | | | | | |
| at AA243598 531 EST #N/A at AA243598 531 EST #N/A at AA252147 551 EST #N/A | | | | | | | | | |
| at AA243598 531 EST #N/A at AA252147 551 EST #N/A | | | | | 42.38 | | 13.66 | | 9.63 |
| at AA252147 551 EST #N/A | | N# | | 60.15 | 62.79 | | | | \ |
| | | /N# | | 95.91 | 56.69 | 122.31 | 3.65 | | |
| EST #N/A | _ | N# | A/N# A | 64.28 | 50.15 | 52.25 | 16.18 | | • |

Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| change sample sample sample sample pormal normal normal lin hcc set 2: set 3: set 3 | | | | fold | | hcc |) Picc | hcc | • | • | - |
|---|-------------------------------------|---------------------------|------|--------|---------|--------|-----------|---------|---------|---------|---------|
| In hate | | | | change | | samble | samble | samble | normal | normal | normal |
| ### #N/A #N/A 184.99 102.43 289.71 8.06 7.61 #N/A #N/A 187.05 63.54 232.2 -4.7 -5.33 #N/A #N/A 137.05 63.54 232.2 -4.7 -5.33 #N/A #N/A 59.19 56.31 23.84 21.4 18.8 #N/A #N/A 524.07 516.6 88.67 441.83 355.04 27.24 0.00498 202.52 192.15 142.78 11.33 18.29 20.72 0.00114 629.56 433.87 551.97 -174.29 -216.08 1 6.09 0.01224 221.56 130.95 237.73 -4.35 -8.44 #N/A #N/A 179.08 84.98 218.93 -1.99 1.49 #N/A #N/A 103.35 97.42 153.31 72.58 70.64 1 #N/A #N/A 258.37 212.57 171.93 133.78 116.63 #N/A #N/A 258.37 212.57 171.93 133.78 116.63 #N/A #N/A 258.37 212.57 171.93 133.78 116.63 #N/A #N/A 32.11 31.27 15.41 8.91 6.74 #16.7 #N/A #N/A 52.26 34.54 55.5 9.04 14.67 #N/A #N/A 91.01 73.62 80.93 24.83 12.83 5.72 0.00384 316.32 322.55 180.22 65.48 28.37 #N/A #N/A 89.98 66.8 71.4 15.89 14.41 \$3.10.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A #N/A 284.91 287.54 145.65 5.00 #N/A #N/A 284.91 287.54 19.03 31.67 73.6 80.63 31.67 547 7.05 #N/A #N/A 39.35 37.8 145.65 50.03 13.67 6.8 71.24 19.03 18.75 #N/A #N/A 39.35 71.21 49.95 87.89 1.67 547.34 #N/A #N/A 39.35 77.31 12.00 31.00 3 | | | | In hcc | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| #N/A #N/A 184.99 102.43 289.71 8.06 7.61 #N/A #N/A 137.05 63.54 232.2 4.7 -5.33 #N/A #N/A 59.19 56.31 23.84 21.4 18.8 #N/A #N/A 524.07 516.6 88.67 441.83 355.04 2 7.24 0.00498 202.52 192.15 142.78 11.33 18.29 20.72 0.00114 629.56 433.87 551.97 -174.29 -216.08 1 6.09 0.01224 221.56 130.95 237.73 4.35 -8.44 #N/A #N/A 179.08 84.98 218.93 -1.99 1.49 #N/A #N/A 200.3 136.72 140.42 118.18 97.38 1 #N/A #N/A 103.35 97.42 153.31 72.58 70.64 1 #N/A #N/A 103.35 97.42 153.31 72.58 70.64 1 #N/A #N/A 128.41 118.8 40.54 65.66 70.33 6.22 0.00001 309.74 308.36 112.62 51.84 43.51 8.75 #N/A #N/A 52.26 34.54 55.5 9.04 14.67 #N/A #N/A 52.26 34.54 55.5 9.04 14.67 #N/A #N/A 91.01 73.62 80.93 24.83 12.83 5.72 0.00384 316.32 322.55 #N/A #N/A 89.98 66.8 71.4 15.89 14.41 #N/A #N/A 89.98 66.8 71.4 15.89 14.41 #N/A #N/A 284.91 287.54 145.66 466.07 547.34 #N/A #N/A 284.91 287.54 145.65 466.07 547.34 #N/A #N/A 284.91 287.54 145.67 54.00 39.35 37.49 45.67 59.03 #N/A #N/A 284.91 287.54 145.67 54.00 39.35 37.49 45.67 57.05 #N/A #N/A 39.35 37.49 45.67 57.05 #N/A #N/A 39.35 37.49 45.67 57.05 31.24 19.03 18.75 | Genbank Seg ID Known Gene Name | ٥ | Name | set 2 | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| #N/A #N/A 137.05 63.54 23.22 -4.7 -5.33 #N/A #N/A 59.19 56.31 23.84 21.4 18.8 #N/A #N/A 524.07 516.6 88.67 441.83 355.04 2 7.24 0.00498 202.52 192.15 142.78 11.33 18.29 20.72 0.00114 629.56 433.87 551.97 -174.29 -216.08 1 6.09 0.01224 221.56 130.95 237.73 -4.35 -8.44 #N/A #N/A 179.08 84.98 218.93 -1.99 1.49 #N/A #N/A 200.3 136.72 140.42 118.18 97.38 1 #N/A #N/A 258.37 212.57 171.93 133.78 116.63 #N/A #N/A 258.37 212.57 171.93 133.78 116.63 #N/A #N/A 258.37 212.57 171.93 133.78 116.63 #N/A #N/A 32.11 31.27 15.41 8.91 6.74 #N/A #N/A 32.11 31.27 15.41 8.91 6.74 #N/A #N/A 52.26 34.54 55.5 9.04 14.67 #N/A #N/A 52.26 34.54 55.5 9.04 14.67 #N/A #N/A 67.33 42.61 65.11 29.14 23.25 #N/A #N/A 89.98 66.8 71.4 15.89 14.41 \$3.10.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A #N/A 39.35 37.49 4.67 54.7 7.05 | AA279667 629 EST | | | #N/A | #N/A | 184.99 | 102.43 | 289.71 | 8.06 | 7.61 | 23.78 |
| #N/A #N/A 59.19 56.31 23.84 21.4 18.8 #N/A #N/A 524.07 516.6 88.67 441.83 355.04 2 7.24 0.00498 202.52 192.15 142.78 11.33 18.29 20.72 0.00114 629.56 433.87 551.97 -174.29 -216.08 1 6.09 0.01224 221.56 130.95 237.73 -4.35 -8.44 #N/A #N/A #N/A 200.3 136.72 140.42 118.18 97.38 1 #N/A #N/A #N/A 103.35 97.42 153.31 72.58 70.64 1 #N/A #N/A #N/A 258.37 212.57 171.93 133.78 116.63 #N/A #N/A #N/A 32.11 31.27 15.41 8.91 6.74 #N/A #N/A #N/A 52.26 34.54 55.5 9.04 14.67 #N/A #N/A #N/A 91.01 73.62 80.93 24.83 12.83 5.72 0.00384 316.32 322.55 180.22 65.48 28.37 #N/A #N/A #N/A 67.33 42.61 65.11 29.14 23.25 #N/A #N/A #N/A 89.98 66.8 71.4 15.89 14.41 \$5.1 0.00329 376.8 34.106 145.28 90 72.97 1 3.1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A #N/A #N/A 284.91 287.54 145.66 466.07 54.7 7.05 #N/A #N/A #N/A 39.35 37.49 4.67 5.47 7.05 #N/A #N/A #N/A 39.35 37.49 4.67 5.47 7.05 | 4A279943 635 EST | | | W/N# | #N/A | 137.05 | 63.54 | | -4.7 | -5.33 | 11.18 |
| #NI/A #NI/A 524.07 516.6 88.67 441.83 355.04 2 7.24 0.00498 202.52 192.15 142.78 11.33 18.29 20.72 0.00114 629.56 433.87 551.97 -174.29 -216.08 1 6.09 0.01224 221.56 130.95 237.73 -4.35 -8.44 #NI/A #NI/A 179.08 84.98 218.93 -1.99 1.49 #NI/A #NI/A 258.37 212.57 171.93 133.78 116.63 #NI/A #NI/A 103.35 97.42 153.31 72.58 70.64 1 #NI/A #NI/A 258.37 212.57 171.93 133.78 116.63 #NI/A #NI/A 258.37 212.57 171.93 133.78 116.63 #NI/A #NI/A 32.11 31.27 15.41 8.91 6.74 #NI/A #NI/A 32.11 31.27 15.41 8.91 6.74 #NI/A #NI/A 32.11 31.27 15.41 8.91 6.74 #NI/A #NI/A 65.26 34.54 55.5 9.04 14.67 #NI/A #NI/A 67.33 42.61 65.11 29.14 23.25 #NI/A #NI/A 89.98 66.8 71.4 15.89 14.41 #NI/A #NI/A 89.98 66.8 71.4 15.89 14.41 #NI/A #NI/A 89.98 66.8 71.4 15.89 17.41 #NI/A #NI/A 284.91 287.54 145.66 466.07 547.34 1 #NI/A #NI/A 284.91 287.54 145.66 466.07 547.34 1 #NI/A #NI/A 39.35 37.49 4.67 5.47 7.05 #NI/A #NI/A 39.35 37.49 19.03 18.75 | AA290594 691 EST | _ | | #N/A | #N/A | 59.19 | 56.31 | | 21.4 | 18.8 | 23.98 |
| 7.24 0.00498 202.52 192.15 142.78 11.33 18.29 20.72 0.00114 629.56 433.87 551.97 -174.29 -216.08 1 6.09 0.01224 221.56 130.95 237.73 -4.35 -8.44 #NI/A #NI/A 179.08 84.98 218.93 -1.99 1.49 #NI/A #NI/A 200.3 136.72 140.42 118.18 97.38 1 #NI/A #NI/A 103.35 97.42 153.31 72.58 70.64 1 #NI/A #NI/A 103.35 97.42 153.31 72.58 70.64 1 #NI/A #NI/A 258.37 212.57 171.93 133.78 116.63 #NI/A #NI/A 32.11 31.27 15.41 8.91 6.74 #NI/A #NI/A 32.11 31.27 15.41 8.91 6.74 #NI/A #NI/A 67.33 42.61 65.11 29.14 23.25 #NI/A #NI/A 67.33 42.61 65.11 29.14 23.25 #NI/A #NI/A 89.98 66.8 71.4 15.89 14.41 5.1 0.00329 376.8 341.06 145.28 90 72.97 1 3.1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #NI/A #NI/A 284.91 287.54 145.66 466.07 547.34 1 #NI/A #NI/A 284.91 287.54 145.66 466.07 547.34 1 #NI/A #NI/A 39.35 37.49 4.67 5.47 7.05 #NI/A #NI/A 181/A 18 | AA290776 693 EST | | | #N/A | #N/A | 524.07 | 516.6 | | 441.83 | 355.04 | 245.84 |
| 20.72 0.00114 629.56 433.87 551.97 -174.29 -216.08 1 6.09 0.01224 221.56 130.95 237.73 -4.35 -8.44 #NI/A #NI/A 179.08 84.98 218.93 -1.99 1.49 #NI/A #NI/A 200.3 136.72 140.42 118.18 97.38 1 #NI/A #NI/A 103.35 97.42 153.31 72.58 70.64 1 #NI/A #NI/A 128.41 118.8 40.54 65.66 70.33 6.22 0.00001 309.74 308.36 112.62 51.84 43.51 #NI/A #NI/A 32.11 31.27 15.41 8.91 6.74 #NI/A #NI/A 32.11 31.27 15.41 8.91 6.74 #NI/A #NI/A 91.01 73.62 80.93 24.83 12.83 5.72 0.00384 316.32 322.55 180.22 65.48 28.37 #NI/A #NI/A 67.33 42.61 65.11 29.14 23.25 #NI/A #NI/A 89.98 66.8 71.4 15.89 14.41 5.1 0.00329 376.8 341.06 145.28 90 72.97 1 3.1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #NI/A #NI/A 284.91 287.54 145.66 466.07 547.34 1 #NI/A #NI/A 284.91 287.54 145.66 466.07 547.34 1 #NI/A #NI/A 39.35 37.49 4.67 5.47 7.05 #NI/A #NI/A 128.75 128.75 131.24 19.03 18.75 | AA292765 712 ZW10 interactor | • | | 7.24 | 0.00498 | 202.52 | 192.15 | | 11.33 | 18.29 | 13.27 |
| 6.09 0.01224 221.56 130.95 237.73 -4.35 -8.44 #NI/A #NI/A 179.08 84.98 218.93 -1.99 1.49 #NI/A #NI/A 200.3 136.72 140.42 118.18 97.38 1 #NI/A #NI/A 103.35 97.42 153.31 72.58 70.64 1 #NI/A #NI/A 128.41 118.8 40.54 65.66 70.33 6.22 0.00001 309.74 308.36 112.62 51.84 43.51 #NI/A #NI/A 32.11 31.27 15.41 8.91 6.74 #NI/A #NI/A 91.01 73.62 80.93 24.83 12.83 5.72 0.00384 316.32 322.55 180.22 65.48 28.37 #NI/A #NI/A 67.33 42.61 65.11 29.14 23.25 #NI/A #NI/A 89.98 66.8 71.4 15.89 14.41 5.1 0.00329 376.8 341.06 145.28 90.72.97 1 5.1 0.00325 71.21 49.95 87.89 -29.4 -30.17 #NI/A #NI/A 284.91 287.54 145.66 466.07 547.34 1 #NI/A #NI/A 284.91 287.54 145.03 18.75 #NI/A #NI/A 39.35 37.49 4.67 5.47 7.05 #NI/A #NI/A 39.35 37.49 14.07 18.75 | AA398908 801 EST | _ | | 20.72 | 0.00114 | 629.56 | 433.87 | | -174.29 | -216.08 | 104.42 |
| #N/A #N/A 179.08 84.98 218.93 -1.99 1.49 #N/A #N/A 80.97 82 30.96 50.41 51.29 #N/A #N/A 200.3 136.72 140.42 118.18 97.38 1 #N/A #N/A 103.35 97.42 153.31 72.58 70.64 1 #N/A #N/A 128.41 118.8 40.54 65.66 70.33 6.22 0.00001 309.74 308.36 112.62 51.84 43.51 #N/A #N/A 32.11 31.27 15.41 8.91 6.74 #N/A #N/A 52.26 34.54 55.5 9.04 14.67 #N/A #N/A 91.01 73.62 80.93 24.83 12.83 5.72 0.00384 316.32 322.55 180.22 65.48 28.37 #N/A #N/A 67.33 42.61 65.11 29.14 23.25 #N/A #N/A 89.98 66.8 71.4 15.89 14.41 5.1 0.00329 376.8 341.06 145.28 90 72.97 1 3.1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A #N/A 39.35 37.49 4.67 5.47 7.05 #N/A #N/A 39.35 37.49 4.67 5.47 7.05 | AA405098 855 EST | _ | | 60.9 | 0.01224 | 221.56 | 130.95 | • | -4.35 | -8.44 | 30.78 |
| #N/A #N/A 200.3 136.72 140.42 118.18 97.38 1 #N/A #N/A 103.35 97.42 153.31 72.58 70.64 1 #N/A #N/A 128.41 118.8 40.54 65.66 70.33 6.22 0.00001 309.74 308.36 112.62 51.84 43.51 #N/A #N/A 32.11 31.27 15.41 8.91 6.74 #N/A #N/A 91.01 73.62 80.93 24.83 12.83 5.72 0.00384 316.32 322.55 180.22 65.48 28.37 #N/A #N/A 67.33 42.61 65.11 29.14 23.25 #N/A #N/A 89.98 66.8 71.4 15.89 14.41 5.1 0.00329 376.8 341.06 145.28 90 72.97 1 3.1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A #N/A 39.35 37.49 4.67 5.47 7.05 #N/A #N/A 10.02 10.00 | AA406542 878 EST | | | A/N# | #N/A | 179.08 | 84.98 | 218.93 | -1.99 | 1.49 | 12.75 |
| #N/A 103.35 97.42 153.31 72.58 70.64 1 #N/A 103.35 97.42 153.31 72.58 70.64 1 #N/A 128.41 118.8 40.54 65.66 70.33 | AA435738 1042 EST | | | #N/A | #N/A | 80.97 | 82 | 30.96 | 50.41 | 51.29 | 33.08 |
| #N/A 103.35 97.42 153.31 72.58 70.64 1 #N/A 258.37 212.57 171.93 133.78 116.63 #N/A 128.41 118.8 40.54 65.66 70.33 0.000001 309.74 308.36 112.62 51.84 43.51 #N/A 32.11 31.27 15.41 8.91 6.74 #N/A 91.01 73.62 80.93 24.83 12.83 0.00384 316.32 322.55 180.22 65.48 28.37 #N/A 67.33 42.61 65.11 29.14 23.25 #N/A 89.98 66.8 71.4 15.89 14.41 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 39.35 37.49 4.67 5.47 7.05 #N/A 73.6 80.62 31.24 19.03 18.75 | AA443585 1077 EST | | | #N/A | #N/A | 200.3 | 136.72 | 140.42 | 118.18 | 97.38 | 102.71 |
| #N/A 103.35 97.42 153.31 72.58 70.64 1 4N/A 258.37 212.57 171.93 133.78 116.63 4N/A 128.41 118.8 40.54 65.66 70.33 0.00001 309.74 308.36 112.62 51.84 43.51 43.51 40.00384 316.32 32.55 180.22 65.48 28.37 40.00384 316.32 322.55 180.22 65.48 28.37 40.00389 376.8 341.06 145.28 90 72.97 10.00329 376.8 341.06 145.28 90 72.97 10.00325 71.21 49.95 87.89 -29.4 -30.17 41.14 28.35 41.06 145.66 466.07 547.34 14.14 15.89 12.83 12 | Ts translation elongation | Ts translation elongation | | • | | | | | | | |
| #N/A 258.37 212.57 171.93 133.78 116.63 #N/A 128.41 118.8 40.54 65.66 70.33 0.000001 309.74 308.36 112.62 51.84 43.51 #N/A 32.11 31.27 15.41 8.91 6.74 #N/A 32.11 31.27 15.41 8.91 6.74 #N/A 91.01 73.62 80.93 24.83 12.83 0.00384 316.32 322.55 180.22 65.48 28.37 #N/A 67.33 42.61 65.11 29.14 23.25 #N/A 89.98 66.8 71.4 15.89 14.41 0.00329 376.8 341.06 145.28 90 72.97 1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 39.35 37.49 4.67 5.47 7.05 #N/A 73.6 80.62 31.24 19.03 18.75 | AA446596 1092 factor, mitochondrial | • | | #N/A | #N/A | 103.35 | 97.42 | 153.31 | 72.58 | | 154.24 |
| #N/A 128.41 118.8 40.54 65.66 70.33 0.00001 309.74 308.36 112.62 51.84 43.51 #N/A 32.11 31.27 15.41 8.91 6.74 #N/A 52.26 34.54 55.5 9.04 14.67 #N/A 91.01 73.62 80.93 24.83 12.83 0.00384 316.32 322.55 180.22 65.48 28.37 #N/A 89.98 66.8 71.4 15.89 14.41 0.00329 376.8 341.06 145.28 90 72.97 1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 73.6 80.62 31.24 19.03 18.75 | | | | #N/A | #N/A | 258.37 | 212.57 | 171.93 | 133.78 | • | 56.34 |
| #N/A 32.11 31.27 15.41 8.91 6.74 #N/A 32.11 31.27 15.41 8.91 6.74 #N/A 52.26 34.54 55.5 9.04 14.67 #N/A 91.01 73.62 80.93 24.83 12.83 0.00384 316.32 322.55 180.22 65.48 28.37 #N/A 67.33 42.61 65.11 29.14 23.25 #N/A 89.98 66.8 71.4 15.89 14.41 0.00329 376.8 341.06 145.28 90 72.97 1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 39.35 37.49 4.67 5.47 7.05 #N/A 73.6 80.62 31.24 19.03 18.75 | | | | #N/A | #N/A | 128.41 | 118.8 | 40.54 | 65.66 | | 40.45 |
| #N/A 32.11 31.27 15.41 8.91 6.74 #N/A 52.26 34.54 55.5 9.04 14.67 #N/A 91.01 73.62 80.93 24.83 12.83 10.00384 316.32 322.55 180.22 65.48 28.37 #N/A 67.33 42.61 65.11 29.14 23.25 #N/A 89.98 66.8 71.4 15.89 14.41 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 18.75 #N/A 73.6 80.62 31.24 19.03 18.75 | AA459254 1211 EST | | | 6.22 | 0.00001 | 309.74 | 308.36 | 112.62 | 51.84 | 43.51 | 37.57 |
| #N/A 32.11 31.27 15.41 8.91 6.74 #N/A 52.26 34.54 55.5 9.04 14.67 #N/A 91.01 73.62 80.93 24.83 12.83 0.00384 316.32 322.55 180.22 65.48 28.37 #N/A 67.33 42.61 65.11 29.14 23.25 #N/A 89.98 66.8 71.4 15.89 14.41 0.00329 376.8 341.06 145.28 90 72.97 1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 73.6 80.62 31.24 19.03 18.75 | chromosome-associated | chromosome-associated | _ | | | | | | | | |
| #N/A 52.26 34.54 55.5 9.04 14.67 #N/A 91.01 73.62 80.93 24.83 12.83 12.83 0.00384 316.32 322.55 180.22 65.48 28.37 #N/A 67.33 42.61 65.11 29.14 23.25 #N/A 89.98 66.8 71.4 15.89 14.41 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 73.6 80.62 31.24 19.03 18.75 | 1220 | | | #N/A | #N/A | 32.11 | 31.27 | 15.41 | 8.91 | 6.74 | 8.86 |
| #N/A 91.01 73.62 80.93 24.83 12.83 10.00384 316.32 322.55 180.22 65.48 28.37 #N/A 67.33 42.61 65.11 29.14 23.25 #N/A 89.98 66.8 71.4 15.89 14.41 0.00329 376.8 341.06 145.28 90 72.97 1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 39.35 37.49 4.67 5.47 7.05 #N/A 73.6 80.62 31.24 19.03 18.75 | AA476333 1282 EST | | | #N/A | #N/A | 52.26 | 34.54 | . 55.5 | 9.04 | 14.67 | 24.04 |
| 0.00384 316.32 322.55 180.22 65.48 28.37 #N/A 67.33 42.61 65.11 29.14 23.25 #N/A 89.98 66.8 71.4 15.89 14.41 0.00329 376.8 341.06 145.28 90 72.97 1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 39.35 37.49 4.67 5.47 7.05 #N/A 73.6 80.62 31.24 19.03 18.75 | AA496993 1405 EST | | | #N/A | #N/A | 91.01 | 73.62 | 80.93 | 24.83 | 12.83 | 29.07 |
| #N/A 67.33 42.61 65.11 29.14 23.25 #N/A 89.98 66.8 71.4 15.89 14.41 0.00329 376.8 341.06 145.28 90 72.97 1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 39.35 37.49 4.67 5.47 7.05 #N/A 73.6 80.62 31.24 19.03 18.75 | AA504512 1415 KIAA0943 protein | | | 5.72 | 0.00384 | 316.32 | 322.55 | 180.22 | 65.48 | 28.37 | 93.94 |
| #N/A 89.98 66.8 71.4 15.89 14.41 1 0.00329 376.8 341.06 145.28 90 72.97 1 1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 39.35 37.49 4.67 5.47 7.05 #N/A 73.6 80.62 31.24 19.03 18.75 | AA609132 1479 EST | | | #N/A | #N/A | 67.33 | 42.61 | 65.11 | 29.14 | 23.25 | 53.79 |
| 1 0.00329 376.8 341.06 145.28 90 72.97 1 1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 39.35 37.49 4.67 5.47 7.05 #N/A 73.6 80.62 31.24 19.03 18.75 | AA609942 1494 EST | | | #N/A | #N/A | 89.98 | 66.8 | 71.4 | 15.89 | 14.41 | 25.58 |
| 1 0.00329 376.8 341.06 145.28 90 72.97 1 1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 39.35 37.49 4.67 5.47 7.05 #N/A 73.6 80.62 31.24 19.03 18.75 | Fas-activated | Fas-activated | | | | | | | | | |
| 1 0.02325 71.21 49.95 87.89 -29.4 -30.17 #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 39.35 37.49 4.67 5.47 7.05 #N/A 73.6 80.62 31.24 19.03 18.75 | F02254 1777 serine/threonine kinase | | | 5.1 | 0.00329 | 376.8 | 341.06 | | 6 | 72.97 | 135.39 |
| #N/A 284.91 287.54 145.66 466.07 547.34 1 #N/A 39.35 37.49 4.67 5.47 7.05 #N/A 73.6 80.62 31.24 19.03 18.75 | H48459 1937 KIAA0186 gene product | | | 3.1 | 0.02325 | 71.21 | 49.95 | | -29.4 | -30.17 | 6.19 |
| #N/A 73.6 80.62 31.24 19.03 18.75 | H87790 2018 EST | | | #N/A | #N/A | 284.91 | 287.54 | • | 466.07 | 547.34 | 192.71 |
| #N/A 73.6 80.62 31.24 19.03 18.75 | N34017 2494 EST | | | #N/A | #N/A | 39.35 | 37.49 | | 5.47 | 7.05 | 10.01 |
| | N35493 2502 EST | _ | | #N/A | #N/A | 73.6 | 80.62 | | 19.03 | 18.75 | 17 |

Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| | | | | fold | | hcc | hcc |)) (| | | |
|------------------|------------|--------|---|--------|--------------|--------|--------|-------------|--------|--------|---------|
| | | | | change | | sample | sample | sample | normal | normal | normal |
| | | | | in hcc | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| Affy ID | Genbank \$ | Sed ID | Known Gene Name | set 2 | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| rc N63604 at | N63604 | 2601 | EST | #N/A | #N/A | 47.18 | 32.72 | 35 | 16.51 | 14.36 | 10.5 |
| rc_N63646_at | N63646 | 2602 | EST | #N/A | #N/A | 479.9 | 367.24 | 240.56 | 182.22 | 137.95 | 6.06 |
| 1 | | | SRY (sex-determining | | | | | | | | |
| rc N69014 s at | N69014 | 2642 | region Y)-box 22 | #N/A | #N/A | 193.21 | 174.83 | 48.23 | 116.34 | 129.47 | 50.74 |
| rc_N69879_s_at | N69879 | 2651 | drebrin 1 | 3.15 | 0.01659 | 88.78 | 84.86 | 61.76 | 3.69 | -2.94 | 21.49 |
| rc N69983 at | N69983 | 2652 | EST | #N/A | #N/A | 93.25 | 96.86 | 49.82 | 76.2 | 93.14 | 54.92 |
| rc N70330 at | N70330 | 2656 | EST | #N/A | #N/A | 108.39 | 69.74 | 105.54 | 33.78 | 27.33 | 16.47 |
| rc_N74018_at | N74018 | 2684 | EST | #N/A | #N/A | 22.65 | 25.34 | 10.26 | 2.34 | 1.47 | 10.41 |
| rc_N89670_at | N89670 | 2709 | EST | #N/A | #N/A | 65.03 | 59.18 | 66.58 | 3.16 | -1.81 | 43.32 |
| rc N93000 at | N93000 | 2727 | EST | #N/A | #N/A | 77.68 | 64.15 | 45.06 | 17.65 | 18.66 | 10.91 |
| rc N99944 s at | N99944 | 2749 | EST | 3.46 | _ | 211.02 | 200.61 | 102.49 | | 50.12 | 29.73 |
| rc R97176 at | R97176 | 3002 | EST | #N/A | | 143.07 | 152.08 | 86.96 | | 58.26 | 38.91 |
| rc_T16226_at | T16226 | 3066 | EST | 7.23 | _ | 323.53 | 312.1 | 209.59 | 33.87 | 22.78 | 38.68 |
| rc_T32108_at | T32108 | 3103 | EST | #N/A | | 180.81 | 165.53 | 68.41 | 148.09 | 152.11 | 92.06 |
| rc W15275 s at | W15275 | 3472 | EST | #N/A | #N/A | 33.62 | 31.78 | 35.23 | 30.09 | 24.85 | 43.25 |
| rc W31906 at | W31906 | 3497 | secretagogin | 6.62 | 0.00926 | 474.89 | 174.52 | 605.68 | 42.41 | 30.95 | 47.65 |
| rc W46286 s at | W46286 | 3526 | EST | 3.68 | 0.00311 | 154.73 | 166.08 | 72.64 | 23.51 | 22.85 | 61.89 |
| ! ! | | | DEAD/H (Asp-Glu-Ala- | | | | | | | | |
| | | | Asp/His) box polypeptide, Y | | | | | | | | |
| rc W60097 at | W60097 | 3556 | chromosome | 4.82 | 4.82 0.04903 | 235.42 | 265.15 | 189.13 | 35.34 | 11.72 | 84.5 |
| l | | | KDEL (Lys-Asp-Glu-Leu) | | | | | | | | |
| rc_W80852_at | W80852 | 3609 | protein retention receptor 3 Wolf-Hirschhorn syndrome | 6.37 | 0.00005 | 184.61 | 159.98 | 100.74 | 11.16 | 11.92 | 36.06 |
| rc AA046745 at | AA046745 | 113 | candidate 1 | 3.33 | | 73.04 | 72.71 | 48.97 | | 12.02 | 7.33 |
| rc_AA052941_at | AA052941 | 121 | EST | 3.36 | 0.00088 | 100.81 | 106.86 | 4 | • | | |
| rc AA058589 f at | AA058589 | 144 | EST | #N/A | W/V# | 44.63 | 51.66 | 34.1 | • | | |
| rc_AA126561 s at | AA126561 | 249 | stanniocalcin | #N/A | #N/A | 51.44 | 44.19 | 31.6 | 7.16 | | 8.55 |
| 1 | | | | | | | | | | | |

Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| change sample sample sain in the case 2: set 3: set | | | | | to <u>ld</u> | | ncc | ncc | 220 | | | |
|--|----|---------------|--------|----------------------------|--------------|---------|--------|--------|---------|--------|--------|---------|
| in hcc set 2: \$\text{set 2} \text{ Frown Gene Name} \text{ set 2} \text{ p value} Mean Metan Me | | | | | change | | sample | sample | sample | normal | normal | normal |
| Seq ID Known Gene Name set 2 p value Mean Mf 313 homolog 4.72 0.00326 398.05 | | , | | | in hcc | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| target of myb1 (chicken) 313 homolog neighbor of A-kinase anchoring protein 95 699 ubiquitin carrier protein 673 EST 741 creatine kinase, brain 742 creatine kinase, brain 744 creatine kinase, brain 745 creatine kinase, brain 746 creatine kinase, brain 747 creatine kinase, brain 748 factor 1 ATP-binding cassette, subfamily F (GCN20), member 972 2 1021 EST v-Ha-ras Harvey rat sarcoma viral oncogene 1075 homolog 1352 enzyme E2L 3 ubiquitin-conjugating 1352 enzyme E2L 3 3.33 0.00211 113.43 1 1457 EST 1448 KIAA0530 protein 1526 EST 1526 EST 1526 EST 1526 EST 1527 SST 1528 EST 1528 EST 1538 EST 1638 EST 1734 integral type I protein 1734 Fixin Harvey 1736 EST 1737 Fixin Harvey 1738 EST 1738 EST 1739 EST 1739 EST 1739 EST 1734 Fixin Harvey 1736 Fixin 1737 Fixin 1734 Fixin 1735 Fixin 1735 Fixin 1 | O | | Seq ID | Known Gene Name | set 2 | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| 313 homolog | | | | target of myb1 (chicken) | | | | | | | | |
| neighbor of A-kinase 326 anchoring protein 95 669 ubiquitin carrier protein 673 EST 741 creatine kinase, brain 742 creatine kinase, brain 647 0.01462 512.64 744 creatine kinase, brain 745 creatine kinase, brain 746 creatine kinase, brain 747 creatine kinase, brain 748 factor 1 748 creatine kinase, brain 749 creatine kinase, brain 741 creatine kinase, brain 647 0.01462 512.64 717.65 718 EST 718 EST 72 #NI/A #NI/A 137.67 73 6.00211 113.43 74 creatine kinase, brain 647 0.01462 512.64 717.67 72 47 0.01462 512.64 717.67 72 47 0.01462 512.64 72 6.021 72 72 72 73 6.00224 240.81 73 6.0224 740.81 740.81 740.81 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.81 740.82 740.82 740.81 740.82 740.82 740.82 740.82 740.82 740.82 740.81 740.82 74 | ⋖ | A146849 | 313 | homolog | 4.72 | 0.00326 | 398.05 | 341.23 | 244.73 | 98.86 | 106.97 | 86.19 |
| 326 anchoring protein 95 8.55 0.00224 240.81 3 669 ubiquitin carrier protein 673 EST 741 creatine kinase, brain 742 creatine kinase, brain 743 cerzyme EZL 3 1280 EST 744 creatine kinase, brain 647 0.01462 512.64 747 creatine kinase, brain 647 0.01462 512.64 748 factor 1 749 creatine kinase, brain 647 0.01462 512.64 741 creatine kinase, brain 647 0.01462 512.64 740 177.68 741 creatine kinase, brain 647 0.01462 512.64 741 creatine kinase, brain 742 factor 1 744 ATP-binding cassette, subfamily F (GCN20), member 745 family F (GCN20), member 746 factor 1 747 EST 748 KIAA0530 protein 748 KIAA0530 protein 748 KIAA0530 protein 758 EST 759 170 1734 760 1734 7734 integral type I protein 774 factor 1 775 EST 776 factor 1 777 EST 777 factor 1 778 EST 778 EST 779 factor 1 779 factor 1 779 factor 1 779 factor 1 770 factor 1 770 factor 1 770 factor 1 770 factor 1 771 factor 1 772 factor 1 773 factor 1 773 factor 1 773 factor 1 774 factor 1 775 factor 1 775 factor 1 776 factor 1 777 factor 1 778 factor 1 778 factor 1 779 factor 1 770 | | | | neighbor of A-kinase | | | | | | | | |
| 669 ubliquitin carrier protein #NI/A #NI/A 382.91 3 673 EST 741 creatine kinase, brain 6.47 0.01462 512.64 1 TNF receptor-associated 840 factor 1 ATP-binding cassette, subfamily F (GCN20), member 972 2 1021 EST v-Ha-ras Harvey rat sarcoma viral oncogene 1075 homolog 1280 EST ubiquitin-conjugating #NI/A #NI/A 61.23 ubiquitin-conjugating #NI/A #NI/A 61.23 ubiquitin-conjugating #NI/A #NI/A 53.83 1450 EST 3.33 0.00211 113.43 1457 EST 3.26 0.00298 78.73 1528 EST #NI/A #NI/A 83.13 1528 EST #NI/A #NI/A 120.05 1538 EST #NI/A #NI/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1736 EST #NI/A #NI/A 150.04 | ⋖ | A149889 | 326 | anchoring protein 95 | 8.55 | 0.00224 | 240.81 | 312.86 | 143.89 | -0.24 | 8.25 | 29.83 |
| 673 EST #N/A #N/A 117.98 1 741 creatine kinase, brain 6.47 0.01462 512.64 1 TNF receptor-associated 840 factor 1 ATP-binding cassette, subfamily F (GCN20), member 672 2 1021 EST #N/A #N/A 190.92 1 1021 EST #N/A #N/A 190.92 1 1280 EST #N/A #N/A 61.23 138.89 1 1280 EST #N/A #N/A 61.23 1420 EST 83.3 0.00211 113.43 1 1457 EST 3.35 0.03215 122.91 152.8 152.8 1 | Q. | A283711 | 699 | ubiquitin carrier protein | #N/A | #N/A | 382.91 | 380.33 | 210.95 | 221.7 | 203.51 | 93.54 |
| TA1 creatine kinase, brain TNF receptor-associated 840 factor 1 ATP-binding cassette, subfamily F (GCN20), member family famil | ۹ | A284153 | 673 | EST | #N/A | #N/A | 117.98 | 127.81 | 72.86 | 53.98 | 43.58 | 32.17 |
| TNF receptor-associated 840 factor 1 ATP-binding cassette, subfamily F (GCN20), member family F (GCN20), family F (GCN20), member family F (GCN20), member family F (GCN20), | ۹ | A335191 | 741 | creatine kinase, brain | 6.47 | 0.01462 | 512.64 | 199.99 | 673.92 | 43.84 | 33.21 | 43.52 |
| 840 factor 1 ATP-binding cassette, subfamily F (GCN20), member family fam | | | | TNF receptor-associated | | | | | | | | |
| ATP-binding cassette, sub- family F (GCN20), member 972 2 1021 EST #N/A #N/A 190.92 1021 EST #N/A #N/A 25.39 104 EST #N/A #N/A 61.23 105 bomolog #N/A #N/A 61.23 106 Ubiquitin-conjugating #N/A #N/A 245.56 1420 EST 3.33 0.00211 113.43 1420 EST 3.33 0.00211 113.43 1448 KIAA0530 protein #N/A #N/A 53.83 1457 EST 3.26 0.0298 78.73 1528 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1735 EST #N/A #N/A 70.05 | _ | AA402642 | 840 | factor 1 | #N/A | #N/A | 137.67 | 59.13 | 211.07 | 35.93 | 19.15 | 31.32 |
| family F (GCN20), member 972 2 1021 EST #N/A #N/A 190.92 1021 EST #N/A #N/A 25.39 v-Ha-ras Harvey rat sarcoma viral oncogene 1075 homolog 3.4 0.00133 138.89 1280 EST #N/A #N/A 61.23 ubiquitin-conjugating #N/A #N/A 245.56 1420 EST 3.33 0.00211 113.43 1420 EST 3.33 0.00211 113.43 1457 EST 3.26 0.00298 78.73 1528 EST 3.26 0.00298 78.73 1528 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1739 EST #N/A #N/A 121.71 1739 EST #N/A #N/A 121.71 | | | | ATP-binding cassette, sub- | | | | ٠ | | | | |
| 972 2 1021 EST #N/A #N/A 190.92 1021 EST #N/A #N/A 25.39 v-Ha-ras Harvey rat 3.4 0.00133 138.89 1280 EST #N/A #N/A 61.23 ubiquitin-conjugating #N/A #N/A 245.56 1352 enzyme E2L 3 #N/A #N/A 245.56 1420 EST 3.33 0.00211 113.43 1448 KIAA0530 protein #N/A #N/A 53.83 1457 EST 3.55 0.03215 122.91 1526 EST 3.26 0.00298 78.73 1528 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1738 EST #N/A #N/A 450.05 4730 EST #N/A #N/A 450.48 | | | | family F (GCN20), member | | | | | | | | |
| 1021 EST #N/A #N/A #N/A 25.39 v-Ha-ras Harvey rat sarcoma viral oncogene 3.4 0.00133 138.89 1075 homolog 3.4 0.00133 138.89 1280 EST #N/A #N/A 61.23 ubiquitin-conjugating #N/A #N/A 245.56 1352 enzyme E2L 3 3.33 0.00211 113.43 1420 EST 3.33 0.00211 113.43 1448 KIAA0530 protein #N/A #N/A 53.83 1457 EST 3.55 0.03215 122.91 1526 EST 3.26 0.00298 78.73 1528 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1738 EST #N/A #N/A 450.05 | ٠. | AA427460 | 972 | 2 | #N/A | #N/A | 190.92 | 187.57 | 78.83 | 92.5 | 83.63 | 49.3 |
| v-Ha-ras Harvey rat sarcoma viral oncogene 1075 homolog 1280 EST ubiquitin-conjugating 1352 enzyme E2L 3 1420 EST 1420 EST 1448 KIAA0530 protein 1457 EST 1526 EST 1520 0.00298 78.73 1528 EST 1528 EST 1734 integral type I protein 1738 EST 1730 EST | • | AA431429 | 1021 | EST | #N/A | #N/A | 25.39 | 21.98 | 9.32 | 5.32 | 4.23 | 4.43 |
| sarcoma viral oncogene 1075 homolog 1280 EST ubiquitin-conjugating 1352 enzyme E2L 3 1420 EST 1448 KIAA0530 protein 1457 EST 1526 EST 1520 0.00298 1520 1 1528 EST 1734 integral type I protein 1738 EST 1730 EST 1730 EST 1730 EST 1730 EST 1731 EST 1731 EST 1732 EST 1733 EST 1734 EST 1735 EST 1735 EST 1736 EST 1737 EST 1737 EST 1738 EST 1738 EST 1739 EST 1739 EST 1730 EST 1730 EST 1730 EST 1731 EST 1731 EST 1732 EST 1733 EST 1734 EST 1735 EST 1736 EST 1737 EST 1737 EST 1738 EST 1738 EST 1739 EST 1730 EST 1730 EST 1730 EST 1731 EST 1731 EST 1732 EST 1733 EST 1734 EST 1735 EST 1735 EST 1736 EST 1737 EST 1737 EST 1738 EST 1738 EST 1739 EST 1730 EST 1740 EST 1750 EST 1 | | | | v-Ha-ras Harvey rat | | | | | | | | |
| 1075 homolog 3.4 0.00133 138.89 1280 EST #N/A #N/A 61.23 ubiquitin-conjugating #N/A #N/A 245.56 1420 EST 3.33 0.00211 113.43 1448 KIAA0530 protein #N/A #N/A 53.83 1457 EST 3.55 0.03215 122.91 1526 EST 3.26 0.00298 78.73 1528 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1738 EST #N/A #N/A 121.71 1739 EST #N/A #N/A 121.71 | | | | sarcoma viral oncogene | | | | | | | | |
| 1280 EST #N/A #N/A 61.23 ubiquitin-conjugating 1352 enzyme E2L 3 #N/A #N/A 245.56 1420 EST 3.33 0.00211 113.43 1448 KIAA0530 protein #N/A #N/A 53.83 1457 EST 3.55 0.03215 122.91 1526 EST 3.26 0.00298 78.73 1528 EST #N/A #N/A 83.13 1573 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1735 EST #N/A #N/A 121.71 1736 EST #N/A #N/A 121.71 | - | AA443316 | 1075 | homolog | 3.4 | 0.00133 | 138.89 | 124.06 | | 40.07 | 41.12 | 32.54 |
| ubiquitin-conjugating 1352 enzyme E2L 3 1420 EST 1448 KIAA0530 protein #N/A #N/A 53.83 1457 EST 1526 EST 1528 EST 1528 EST 153 EST 1734 integral type I protein 3.17 0.04549 153.76 1738 EST 1730 EST 1731 EST 1731 EST 1732 EST 1733 EST 1734 MN/A #N/A 121.71 | • | AA476260 | 1280 | EST | #N/A | #N/A | 61.23 | 69.49 | 40.45 | -32.66 | -15.85 | 56.91 |
| 1352 enzyme E2L 3 #N/A #N/A 245.56 1420 EST 3.33 0.00211 113.43 1448 KIAA0530 protein #N/A #N/A 53.83 1457 EST 3.55 0.03215 122.91 1526 EST 3.26 0.00298 78.73 1528 EST #N/A #N/A 83.13 1573 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1736 EST #N/A #N/A 70.05 1737 EST #N/A #N/A 150.18 | | | | ubiquitin-conjugating | | | | | | | | |
| 1420 EST 3.33 0.00211 113.43 1448 KIAA0530 protein #N/A #N/A 53.83 1457 EST 3.55 0.03215 122.91 1526 EST 3.26 0.00298 78.73 1528 EST #N/A #N/A 83.13 1573 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1738 EST #N/A #N/A 70.05 1739 EST #N/A #N/A 150.18 | | AA487058 | 1352 | enzyme E2L 3 | #N/A | #N/A | 245.56 | 225.48 | 137.4 | 315.7 | 232.85 | 213.47 |
| 1448 KIAA0530 protein #N/A #N/A 53.83 1457 EST 3.55 0.03215 122.91 1526 EST 3.26 0.00298 78.73 1528 EST #N/A #N/A 83.13 1573 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1738 EST #N/A #N/A 150.05 | | AA521149 | 1420 | EST | 3.33 | 0.00211 | 113.43 | 106.74 | 60.79 | 28.44 | 29.54 | 18.61 |
| 1457 EST 3.55 0.03215 122.91 1526 EST 3.26 0.00298 78.73 1528 EST #N/A #N/A 121.71 1573 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1738 EST #N/A #N/A 70.05 1739 EST #N/A #N/A 150.18 1 | | AA599244 | 1448 | KIAA0530 protein | #N/A | #N/A | 53.83 | 48.84 | 25.7 | 7.71 | 90.9 | 7.11 |
| 1526 EST 3.26 0.00298 78.73 1528 EST #N/A #N/A 121.71 1573 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1738 EST #N/A #N/A 70.05 1739 EST #N/A #N/A 150.18 1 | | AA599850 | 1457 | EST | 3.55 | 0.03215 | 122.91 | 75.32 | 123.52 | -1.23 | -2.33 | 31.86 |
| 1528 EST #N/A #N/A 83.13 1573 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1738 EST #N/A #N/A 70.05 | | AA621530 | 1526 | EST | 3.26 | 0.00298 | 78.73 | 75.43 | 40.45 | 12.46 | 15.12 | 13.43 |
| 1573 EST #N/A #N/A 121.71 1734 integral type I protein 3.17 0.04549 153.76 1738 EST #N/A #N/A 70.05 | | AA621644 | 1528 | EST | #N/A | #N/A | 83.13 | 88.29 | 43.19 | 71.19 | 66.84 | 29.87 |
| 1734 integral type I protein 3.17 0.04549 153.76 1738 EST #N/A #N/A 7730 EST #N/A #N/A 150.05 | | C15078 | 1573 | EST | #N/A | #N/A | 121.71 | 98.25 | 100.16 | 26.22 | 31.14 | 60.2 |
| 1738 EST #N/A #N/A 70.05 | | D80710 | 1734 | integral type I protein | 3.17 | 0.04549 | 153.76 | 74.74 | 166.28 | 22.95 | 28.29 | 24.59 |
| 1730 EST #NIA #NIA 15018 | | D80948 | 1738 | EST | #N/A | #N/A | 70.05 | 77.71 | 32.11 | 41.25 | 43.53 | 17.46 |
| 01.001 ANH ANH | | D81048 | 1739 | EST | W/V# | #N/A | 150.18 | 149.32 | 51.77 | 86.08 | 85.08 | 39.11 |

Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| | | | | ploj | |)) | hcc | hcc | | 1 | - |
|-----------------------|----------------|--------|---------------------------------------|--------|---------|--------|--------|---------|---------|---------|---------|
| | | | | change | | samble | samble | samble | погта | погшан | погшан |
| | | | | in hcc | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| Affy ID | Genbank Seq ID | Sed ID | Known Gene Name | set 2 | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | replication factor C | | | | | | | | |
| rc F04320 s at | F04320 | 1786 | (activator 1) 4 (37kD) | 6.29 | 0.00042 | 156.76 | 122.53 | 109.02 | 14.92 | 17.26 | 9.79 |
| rc H14617 at | H14617 | 1884 | EST | #N/A | #N/A | 143.86 | 147.93 | 66.03 | 133.9 | 150.35 | 53.38 |
| rc_H26763_at | H26763 | 1907 | EST | #N/A | #N/A | 408.05 | 321.65 | 219.8 | 409.85 | 391.85 | 155.84 |
| rc_H68794_at | H68794 | 1984 | EST | 3.67 | 0.00327 | 215.75 | 176.78 | 110.42 | 65.56 | 54 | 51.09 |
| rc H78211 at | H78211 | 2001 | EST | 7.5 | 0.02674 | 285.43 | 307.06 | 261.02 | -115.95 | -129.18 | 70.22 |
| rc H95566 f at | H95566 | 2050 | EST | #N/A | #N/A | 136.58 | 143.31 | 26.27 | 74.11 | 89.04 | 126.61 |
| rc H99870 at | H99870 | 2082 | EST | #N/A | #N/A | 92.81 | 94.93 | 33.98 | 53.96 | 61.17 | 29.86 |
| rc_N62487_at | N62487 | 2591 | EST | #N/A | #N/A | 56.6 | 52.26 | 33.75 | 13.89 | 16.02 | 8.53 |
| | N67815 | 2627 | EST | 3.84 | _ | 351.55 | 356.17 | 99.35 | 104.07 | 151.77 | 86.01 |
| rc N69084 i at | N69084 | 2643 | EST | 3.11 | | 233.32 | 208.16 | 169.31 | 72.13 | 2.79 | 49.99 |
| rc N73278 at | N73278 | 2672 | EST | #N/A | | 77.72 | 71.59 | 49.4 | 25.36 | 19.22 | 26.85 |
| rc R06251 f at | R06251 | 2764 | tumor protein D52-like 2 | 4.88 | 0.03097 | 466.88 | 341.39 | 380.43 | 74.25 | 66.72 | 64.11 |
|]] | | | carbohydrate (keratan | | | | | | | | |
| | | | sulfate Gal-6) | | | | | | | | |
| rc R15740 at | R15740 | 2791 | sulfotransferase 1 | 3.36 | 0.00268 | 117.43 | 100.03 | 64.73 | 31.77 | 21.3 | |
| rc_R24507_at | R24507 | 2802 | EST | #N/A | #N/A | 31.61 | 16.41 | 30.69 | က | 3.45 | |
| rc <u>R27296</u> f at | R27296 | 2807 | EST | 3.41 | 0.00309 | 77.08 | 67.51 | 39.42 | 8.27 | 11.07 | |
| | R30931 | 2810 | EST | #N/A | #N/A | 54.19 | 57.09 | 29.34 | 17.58 | 13.67 | 14.21 |
| rc R45994 f at | R45994 | 2867 | | 6.48 | 0.00358 | 730.52 | 598.82 | 420.14 | 145.88 | 160.69 | 163.03 |
| rc R70253 at | R70253 | 2945 | | 3.38 | 0.03125 | 99.7 | 88.01 | 87.93 | | | |
| rc R72886 s at | R72886 | 2956 | | 5.5 | 0.00091 | 564.05 | 473.4 | 207.71 | 126.66 | 121.18 | _ |
| rc R96924 s at | R96924 | 3001 | EST | 7.04 | 0.00012 | 336.8 | 329 | 106.3 | | ٠, | 54.09 |
| rc_T15852_f_at | T15852 | 3062 | EST | 5.21 | 0.00642 | 168.18 | 134.42 | 120.18 | -7.59 | 7.36 | |
| 1 | | | cleavage and polyadenylation specific | | | | | | | | |
| rc_T16983_s_at | T16983 | 3074 | factor 4, 30kD subunit | 4.23 | 0.0106 | 262.26 | 268.2 | 155.86 | 65.64 | 45.67 | 74.28 |
| | | | | | | | | | | | |

Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| | | | | fold | | hcc | hcc | pcc | | | |
|----------------|-------------|--------|----------------------------|--------|---------|--------|--------|---------|--------|--------|---------|
| | | | | change | | sample | sample | sample | normai | normal | normal |
| | | | | In hcc | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| Affy ID | Genbank Seq | Sed ID | Known Gene Name | set 2 | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | RNA (guanine-7-) | | | | | | | | |
| rc_T33865_f_at | T33865 | 3110 | methyltransferase | #N/A | W/A | 29.73 | 32.86 | 22.11 | -0.26 | 2.33 | 8.15 |
| rc_T66935_at | T66935 | 3179 | EST | 3.01 | 0.00123 | 181.34 | 189.67 | 61.49 | 8.99 | 52.94 | 42.32 |
| rc_T95057_f_at | T95057 | 3259 | EST · | 6.46 | 0.00613 | 229.11 | 188.71 | 174.46 | 7.13 | 15.37 | 37.2 |
| rc_W45320_f_at | W45320 | 3523 | KRAB-associated protein 1 | 10.05 | 0.00002 | 365.22 | 345.46 | 124.13 | -2.52 | 25.03 | 112.32 |
| AF006041_at | AF006041 | 1549 | death-associated protein 6 | #N/A | #N/A | 174.22 | 170.16 | 37.64 | 74.91 | 58.55 | 50.31 |
| D00596_at | D00596 | 1590 | thymidylate synthetase | 5.58 | 0.0098 | 200.17 | 128.11 | 170.5 | 20.16 | 20.43 | 12.43 |
| D38491_at | D38491 | 1653 | KIAA0117 protein | #N/A | #N/A | 47.67 | 52.03 | 21.01 | 20.77 | 20.88 | 17.43 |
| D63486_at | D63486 | 1712 | KIAA0152 gene product | 3.56 | 0.00063 | 277.42 | 248.24 | 99.25 | 84.6 | 95.61 | 45.48 |
| | | | minichromosome | | | | | - | | | |
| | | | maintenance deficient | | | | | | | | |
| D84557_at | D84557 | 1749 | (mis5, S. pombe) 6 | 3.97 | 0.0017 | 184.48 | 132.11 | 125.74 | 37.22 | 43.12 | 28.15 |
| D86957_at | D86957 | 1754 | KIAA0202 protein | 3.08 | 0.02949 | 91.22 | 73.14 | 89.88 | 14.7 | 12.7 | 6.29 |
| D86972_at | D86972 | 1755 | KIAA0218 gene product | #N/A | | 68.56 | 66.54 | 32.32 | 34.58 | 32.05 | 21.82 |
| | | | amylase, alpha 2B; | | | | | | | | |
| D90097_at | D90097 | 1768 | pancreatic | #N/A | #N/A | 41.19 | 37.49 | 21.85 | 6.55 | 11.44 | 15.06 |
| L29218_at | L29218 | 2190 | CDC-like kinase 2 | 3.82 | 0.00035 | 155.42 | 142.58 | 76.28 | 35.11 | 41.27 | 26.41 |
| | | | RNA-binding protein | | | | • | | | | |
| L38696_at | T38696 | 2208 | (autoantigenic) | #N/A | #N/A | 127.32 | 114.91 | 62.34 | 60.18 | 58.55 | 24.77 |
| M11749_at | M11749 | 2240 | Thy-1 cell surface antigen | #N/A | #N/A | 83.1 | 65.86 | 64.52 | 26.05 | 28.95 | 20.56 |
| M25753_at | M25753 | 2308 | cyclin B1 | #N/A | #N/A | 44.14 | 30.96 | 40.36 | -0.8 | 3.03 | 15.35 |
| | | | intercellular adhesion | | | | | | | | |
| M32334_at | M32334 | 2333 | molecule 2 | #N/A | #N/A | 95.04 | 92.33 | 67.51 | 42.19 | 41.31 | 27.33 |
| | | | laminin, gamma 1 (formerly | | | | | | | | |
| M55210_at | M55210 | 2353 | LAMB2) | 3.47 | 0.02551 | 137.63 | 115.85 | 102.71 | 27.07 | 29.17 | 12.36 |
| M61916_at | M61916 | 2372 | laminin, beta 1 | 3.18 | 0.01171 | 75.1 | 67.4 | 61.41 | 7.19 | 7.98 | 8.99 |

Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| | | • | | | | | | | _ | | _ | | ١. | _ | ~ | | ٥, | | <u>ر</u> | | | <u></u> | • | (0 | | " | œ |
|------|----------|--------|-----------------|-----------------|------------------|-------------------------|--------------|----------------------|------------------------|--------------------------|---------------------|-----------------------------|---------------|-------------------------|------------------|-----------------------|-----------|-----------------------------|-----------|----------------------|-----------------------------|---------------|-------------------------|----------------------------|------------------------------|----------------|---------------------|
| | normal | set 2: | Std Dev | | | | 39.03 | | 16.79 | | 101.89 | | 7.55 | 26.9 | 9.38 | | 10.2 | | 23.02 | | | 19.73 | | 11.96 | | 17.66 | 11.78 |
| | normal | set 2: | Median | | | 1 | 20.63 | | 9.92 | | -175.43 | | 12.85 | -1.2 | 7.6 | | 17.43 | | 36.29 | , | | -1.83 | | 14.81 | | 45.39 | 3.77 |
| | normal | set 2: | Mean | | | | 8.67 | | 4.26 | | -155.4 | | 11.53 | 3.36 | 6.3 | | 15.7 | | 36.35 | | | 3.1 | | 16.58 | | 45.63 | 4.73 |
| hcc | sample | set 2: | Std Dev | | | | 192.98 | | 79.29 | | 1414.43 | | 18.34 | 109.13 | 21.66 | | 66.3 | | 31.81 | | | 70.13 | | 61.79 | | 24.25 | 36.93 |
| hcc | sample | set 2: | Median S | | | | 248.21 | | 82.61 | | 193.74 | | 29.21 | 115.3 | 26.91 | | 70.23 | | 61.64 | | | 98.2 | | 83.66 | | 93.3 | 48.48 |
| hcc | sample s | set 2: | Mean | ! | | | 216.51 | | 110.39 | | 690.32 | | 29.33 | 143.52 | 25.79 | | 83.43 | | 73.85 | | | 84.5 | | 97.76 | | 84.78 | 56.15 |
| | •, | | p value | | | | 5.15 0.02881 | | 0.00116 | | 0.02104 | | #N/A | 0.00547 | #N/A | | #N/A | | #N/A | | | 0.02469 | | 0.0045 | | #N/A | #N/A |
| fold | change | in hcc | set 2 | | | | 5.15 | | 4.59 | | 9.86 | | #N/A | 4.83 | #N/A | | #N/A | | #N/A | | | 3.12 | | 3.55 | | #N/A | #N/A |
| | | | Known Gene Name | stress-induced- | phosphoprotein 1 | (Hsp70/Hsp90-organizing | protein) | replication factor C | (activator 1) 4 (37kD) | midkine (neurite growth- | promoting factor 2) | cytosolic ovarian carcinoma | antigen 1 | cell division cycle 25B | laminin, alpha 4 | TAR (HIV) RNA-binding | protein 1 | diacylglycerol kinase, zeta | (104kD) | DEAD/H (Asp-Glu-Ala- | Asp/His) box polypeptide 17 | (72kD) | gamma-aminohutvric acid | (GABA) A recentor, ensilon | multiple endocrine neoplasia | | bleomycin hydrolase |
| | | | Seq ID | | | | 2412 | | 2415 | | 2426 | | 3031 | 3036 | 3037 | | 3357 | | 3384 | | | 3398 | | 3406 | | 3462 | 3822 |
| | | | Genbank Seq ID | | | | M86752 | | M87339 | | M94250 | | S72904 | S78187 | S78569 | | U38847 | | U51477 | | | U59321 | | 1166661 | | U93237 | X92106 |
| | | | Affy ID | | | , | M86752_at | | M87339_at | | M94250_at | | S72904 at | S78187 at | S78569 at | l | U38847_at | | U51477_at | | | U59321_at | | 1166661 at | 5 | U93237 rna2 at | X92106_at |

Table 8A. Genes and ESTs expressed only in hepatocellular carcinoma versus normal sample set 2.

| Affy ID Genbank Seq ID Known Gene Name tafazzin (cardiomyopathy, dilated 3A (X-linked); endocardial fibroelastosis 2; S2762 3828 Barth syndrome) dual-specificity tyrosine-(Y)-phosphorylation regulated Y09216 3856 kinase 2 guanine nucleotide binding protein (G protein), alpha 11 M69013_at M69013 2391 (Gq class) AD000092_cds7_s_at AD000092 1542 homolog A small nuclear ribonucleoprotein 70kD x04654_s_at X04654 3681 polypeptide (RNP antigen) myosin, light polypeptide 1, myosin, light polypeptide 1, alpha 1205648_s_at W74715_s_at M74715 2395 duronidase, alpha-L-a disintegrin and metalloproteinase domain U41767_s_at W12125_at M12125_at tropomyosin 2 (beta) sulfortransferase family 1A, sulforted 1A, sulfortransferase family 1A, sulfortran | | fold | • | hcc | hcc | hcc | | | |
|--|--------------------------------|--------|----------|--------|--------|---------|--------|--------|---------|
| Seq ID 3828 3828 2391 1542 2326 2326 3379 2395 | | cnange | <i>,</i> | sample | samble | sample | погла | погта | погша |
| 3828 3828 3856 1542 1542 3681 2326 2336 3360 2241 | | In hcc | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| 3828 3856 2391 1542 3681 3379 2395 3360 2241 | Seq ID | set 2 | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| 3828 3856 2391 1542 3379 2395 3360 2241 | tafazzin (cardiomyopathy, | | | | | | | | |
| 3828 3856 2391 1542 3379 2395 2395 3360 2241 | dilated 3A (X-linked); | | | | | | | | |
| 3828 3856 2391 1542 3379 2326 2395 3360 2241 | endocardial fibroelastosis 2; | | | | | | | | |
| 3856 2391 1542 3681 2326 2395 3360 2241 | | #N/A | #N/A | 82.96 | 83.94 | 15.05 | 23.4 | 33.5 | 24.15 |
| 3856 2391 1542 3681 2326 2395 3360 2241 | dual-specificity tyrosine-(Y)- | | | | | | | | |
| 3856 2391 1542 3379 2326 2395 3360 2241 | phosphorylation regulated | | | | | | | | |
| 2391 1542 3681 2326 2395 2395 2395 | _ | #N/A | #N/A | 47.87 | 47.61 | 18.08 | 18.99 | 22.48 | 13.31 |
| 2391 1542 3681 2326 2395 2395 3360 2241 | guanine nucleotide binding | | | | | | | | |
| 2391 1542 3681 2326 2395 2395 3360 2241 | protein (G protein), alpha 11 | - | | | | | | | |
| 3681 2326 2326 2395 3379 3360 2241 | | #N/A | #N/A | 86.89 | 70.61 | 47.65 | 38.18 | 47.33 | 23.06 |
| 3681 2326 2326 2395 3360 2241 | | | | | | | | | |
| 3681 2326 3379 2395 3360 2241 | 1542 | #N/A | #N/A | 97.12 | 98.23 | 25.66 | 37.18 | 32.35 | 28.04 |
| X04654 3681 M31211 2326 U50648 3379 M74715 2395 U41767 3360 M12125 2241 | | | | | | | | | |
| X04654 3681 M31211 2326 U50648 3379 M74715 2395 U41767 3360 M12125 2241 | ribonucleoprotein 70kD | | | | | | | | |
| M31211 2326 U50648 3379 M74715 2395 U41767 3360 M12125 2241 | 3681 polypeptide (RNP antigen) | #N/A | #N/A | 122.45 | 134.9 | 49.51 | 42.22 | 41.24 | 17.18 |
| M31211 2326 U50648 3379 M74715 2395 U41767 3360 M12125 2241 | myosin, light polypeptide 1, | | | | | | | | |
| U50648 3379 M74715 2395 U41767 3360 M12125 2241 | | #N/A | #N/A | 60.22 | 49.75 | 54.83 | -2.19 | -4.38 | 11.56 |
| U50648 3379 M74715 2395 U41767 3360 M12125 2241 | protein kinase, interferon- | | | | | | | | |
| U50648 3379 M74715 2395 U41767 3360 M12125 2241 | inducible double stranded | | | | | | | | |
| M74715 2395 U41767 3360 M12125 2241 | _ | #N/A | #N/A | 214.34 | 192.86 | 122.38 | 105.38 | 92.58 | 43.12 |
| U41767 3360 M12125 2241 | _ | #N/A | #N/A | 122.69 | 121.95 | 72.58 | 41.88 | 24.48 | 43.99 |
| U41767 3360 M12125 2241 | a disintegrin and | | | | | | | | |
| U41767 3360 M12125 2241 | metalloproteinase domain | | | | | | | | |
| M12125 2241 | • | W/V# | #N/A | 196.76 | 147.11 | 113.93 | 83.03 | 66.93 | 68.33 |
| | • | 7.13 | 0.0004 | 174.25 | 183.97 | 95.24 | 13.01 | 16.33 | 11.64 |
| | sulfotransferase family 1A, | | | | | ٠ | | | |
| | pnenoi-preferring, member | | | | | | | | |
| U20499_at U20499 3321 3 | 3321 3 | W/N# | #N/A | 141.35 | 126.33 | 135.57 | 48.34 | 54.37 | 24.69 |

Table 8B. Genes and ESTs expressed only in normal liver compared to hepatocellular carcinoma.

| | | | | | | 32 | ည္ဌ | hcc | | | |
|------------------|----------|--------|-------------------------------------|----------|---------|--------|--------|---------|--------|--------|---------|
| | | | | | | sample | sample | sample | normal | normal | normal |
| | | | | hcc fold | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | change | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| rc_AA010205_at | AA010205 | 23 | EST | 5.71 | 0.00014 | 26.92 | 20.06 | 32.19 | 187.55 | 154.99 | 92.13 |
| | | • | potassium voltage-gated | | | | | | | | |
| rc_AA013095_s_at | AA013095 | 33 | channel, shaker-related | #N/A | #N/A | 5.12 | 7.9 | 12.98 | 18.56 | 15.83 | 8.24 |
| rc_AA046747_at | AA046747 | 114 | EST | 4.77 | 0.00023 | -0.57 | 3.68 | 23.81 | 113.78 | 88.54 | 66.41 |
| rc_AA074885_at | AA074885 | 161 | macrophage receptor with | 10.88 | 0.00087 | 35.67 | 58.01 | 44.34 | 652.03 | 761.74 | 300.57 |
| rc_AA099225_at | AA099225 | 206 | EST | 6.59 | 0.00064 | 10.9 | 9.75 | 15.11 | 212.68 | 163.45 | 194.31 |
| rc_AA253410_at | AA253410 | 564 | EST | #N/A | #N/A | 3.66 | 1.48 | 10.67 | 49.37 | 26.82 | 45.2 |
| rc_AA281796_at | AA281796 | 650 | mannose-P-dolichol utilitzation | #N/A | #N/A | 95.74 | 105.86 | 45.46 | 170.88 | 165.02 | 41.87 |
| rc_AA282541_at | AA282541 | 661 | EST | #N/A | #N/A | 17.21 | 11.75 | 13.38 | 31.31 | 29.9 | 13.53 |
| rc_AA285053_at | AA285053 | 681 | EST | ß | 0.00718 | 36.49 | 20.15 | 49.24 | 238.16 | 242.27 | 169.12 |
| rc_AA287566_at | AA287566 | 069 | KIAA0187 gene product | 6.9 | 0.00023 | 18.67 | 14.05 | 35.71 | 246.24 | 201.66 | 228.64 |
| rc_AA404248_at | AA404248 | 847 | EST | #N/A | #N/A | 24.05 | 30.59 | 15.83 | 40.92 | 35.06 | 17.49 |
| rc_AA448002_at | AA448002 | 1113 | putative type II membrane | 14.14 | 0 | 39.9 | 38.99 | 13.33 | 594.13 | 528.63 | 282.58 |
| rc_AA460661_at | AA460661 | 1229 | EST | 5.46 | 0.00151 | 20.05 | 8.57 | 39.04 | 184.62 | 198.21 | 108.17 |
| rc_AA487576_at | AA487576 | 1357 | EST | #N/A | #N/A | 11.67 | 2.42 | 27.75 | 26.05 | 25.27 | 11.13 |
| rc_AA621235_at | AA621235 | 1517 | EST | #N/A | #N/A | 62.79 | 68.63 | 35.33 | 114.75 | 113.36 | 65.35 |
| rc_F09979_at | F09979 | 1809 | EST . | #N/A | W/A | 61.21 | 6.31 | 128.69 | 226.47 | 116.41 | 288.21 |
| | | | butyrobetaine (gamma), 2- | | | | | | | | |
| rc_F10380_at | F10380 | 1816 | oxoglutarate dioxygenase | #N/A | #N/A | 43.64 | 47.09 | 18.15 | 106.49 | 93.71 | 83.35 |
| rc_H57056_at | H57056 | 1953 | EST | #N/A | #N/A | -1.79 | -7.37 | 11.38 | 35.07 | 38.32 | 17.88 |
| rc_H58673_at | H58673 | 1959 | EST | 15.49 | 0.00002 | 34.96 | 26.37 | 38.53 | 652.47 | 677.55 | 376.36 |
| rc_H59136_at | H59136 | 1962 | EST | 6.63 | 0.00033 | 33.12 | 21.17 | 42.92 | 250.23 | 229.94 | 129.12 |
| H66367_at | H66367 | 1977 | EST | 3.84 | 0.00133 | 37.14 | 35.15 | 26.61 | 155.38 | 149.31 | 70.77 |
| H72650_at | H72650 | 1994 | EST | #N/A | #N/A | 78.97 | 93.74 | 40.9 | 122 | 125 | 42.24 |
| rc_H87765_at | H87765 | 2017 | KIAA0626 gene product | 3.94 | 0.00123 | 10.11 | 10.07 | 4.23 | 94.26 | 93.78 | 60.64 |
| M10943_at | M10943 | 2234 | metallothionein 1F (functional) | 6.23 | 0.00007 | 35.67 | 28.76 | 22.4 | 217.65 | 186.71 | 86.73 |
| M30185_at | M30185 | 2321 | cholesteryl ester transfer protein, | 3.82 | 0.00131 | -15.66 | -12.63 | 23.86 | 93.07 | 86.04 | 61.35 |
| M63967_at | M63967 | 2378 | aldehyde dehydrogenase 5 | 3.88 | 0.00274 | 40.43 | 30.04 | 29.05 | 164.38 | 178.74 | 87.39 |

Table 8B. Genes and ESTs expressed only in normal liver compared to hepatocellular carcinoma.

| | | | | | | hcc | hcc | hcc | | | |
|----------------|----------------|--------|----------------------------------|----------|---------|--------|--------|---------------|---------|---------|---------|
| | | | | | | sample | sample | sample | normal | normal | normal |
| | | | · | hcc fold | | set 2: | set 2: | set 2: | set 2: | set 2: | set 2: |
| Affy ID | Genbank Seq ID | Seq ID | Known Gene Name | change | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| rc_N22404_at | N22404 | 2450 | EST | 3.44 | 0.02267 | 56.69 | 35.82 | 73.26 | 193.63 | 130.09 | 162.83 |
| rc_N34919_at | N34919 | 2499 | EST | #N/A | #N/A | 19.32 | 19.72 | 14.37 | 52.96 | 44.23 | 24.23 |
| rc_N54604_at | N54604 | 2570 | EST | #N/A | #N/A | 45.27 | 14.34 | 81.45 | 133.06 | 119.1 | 109.95 |
| rc_N65959_at | N65959 | 2612 | EST | 3.38 | 0.00785 | 37.8 | 28.94 | 31.45 | 142.87 | 149.26 | 79.26 |
| rc_N67105_at | N67105 | 2624 | EST | 4.69 | 0.00194 | 12.46 | 10.45 | 12.64 | 143.36 | 102.9 | 121.4 |
| rc_N69114_at | N69114 | 2644 | nuclear receptor subfamily 1, | #N/A | #N/A | 12.69 | 5.95 | 15.01 | 72.29 | 73.28 | 57.34 |
| rc_N74624_at | N74624 | 2688 | collectin sub-family member 10 | #N/A | #N/A | 39.43 | 35.17 | 24.21 | 71.81 | 65.09 | 43.57 |
| N99542_at | N99542 | 2747 | orosomucoid 1 | 3.53 | 0.00607 | 61.09 | 67.97 | 39.3 | 226.06 | 187.68 | 146.11 |
| rc_R09053_at | R09053 | 2783 | EST | 3.45 | 0.03074 | 64.12 | 36.33 | 102.03 | 186.87 | 204.61 | 113.19 |
| rc_R73816_at | R73816 | 2961 | EST | 7.05 | 0.01287 | 35.12 | 13.76 | 100.6 | 462.31 | 374.67 | 450.91 |
| | | | CD5 antigen-like (scavenger | | | | | | | | |
| rc_R99591_at | R99591 | 3016 | receptor cysteine rich family) | 7.41 | 0.00043 | 124.14 | 139.76 | 50.35 | 1035.39 | 904.92 | 691.83 |
| rc_T58756_at | T58756 | 3156 | EST | 16.61 | 0 | -1.02 | 1.06 | 24.36 | 390.04 | 386.76 | 164.41 |
| T68083_at | T68083 | 3185 | short-chain | #N/A | #N/A | 96.34 | 59.71 | 104.53 | 263.22 | 280.22 | 136.48 |
| rc_T68711_at | T68711 | 3188 | EST | 35.98 | 0.0003 | 29.03 | -33.95 | 176.15 | 1388.67 | 1074.67 | 890.91 |
| rc_W48860_at | W48860 | 3537 | EST | #N/A | #N/A | 37.25 | 31.7 | 27.5 | 52.69 | 41.31 | 21.1 |
| | | | X-prolyl aminopeptidase | • | | | | | | | |
| rc_W78093_at | W78093 | 3601 | (aminopeptidase P) 2, | W/A | #N/A | 408.58 | 397.88 | 102.27 | 518.77 | 518.8 | 211.32 |
| rc_W85890_at | W85890 | 3623 | EST | #N/A | #N/A | 60.91 | 60.47 | 23.73 | 70.98 | 66.52 | 24.66 |
| rc_W90018_f_at | W90018 | | EST | #N/A | #N/A | 21.51 | 1.31 | 56.9 | 59.32 | 53.19 | 41.68 |
| X54380_at | X54380 | 3730 | pregnancy-zone protein | 7.71 | 0.00069 | 15.11 | 8.07 | 27.07 | 274.41 | 255.4 | 203.47 |
| Z48475_at | 248475 | 3943 | glucokinase (hexokinase 4) | 4.6 | 0.01693 | 62.99 | 51.37 | 146.72 | 305.23 | 239.52 | 155.98 |
| Z49269 at | 749269 | 3945 | subtamily A (Cys-Cys), member 14 | 7.24 | 0.01047 | 138 63 | 66 95 | 201 93 526 13 | 526 13 | 532 29 | 166.67 |
| | | | | | | | 20:55 | 201:02 | 250:10 | 202.20 | 0.00 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | fold | | | | | normal | normal | normal |
|------------------|----------|--------|--|----------|---------|-------------|-------------|-------------------------------------|--------------------|--------|---------|
| | | | | change | - | netastatic: | metastatic: | metastatic: metastatic: metastatic: | set 2: | set 2: | set 2: |
| Affv ID | Genbank | Sed ID | Known Gene Name | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| rc AA007160 at | | 16 | | 9 | 0.01035 | 171.32 | 175.07 | 138.03 | 9.1 | 8.6 | 12.54 |
| 1 | | | diptheria toxin | | | | | | | | |
| | | | resistance protein | | | | | | | | |
| | | | required for | | | | | | | | |
| | | | diphthamide | | | | | | | | |
| | | | biosynthesis | | | | | | | | |
| | | | (Saccharomyces)-like | | | | | | | | |
| AA009913 at | AA009913 | 21 | 2 | #N/A | #N/A | 69.53 | 81.03 | 27.69 | 16.81 | 18.22 | 13.51 |
| rc AA011134 at | AA011134 | 59 | EST | 28.79 | 0.00602 | 1157.47 | 566.78 | 1448.16 | -72.57 | -51.2 | 69.3 |
|] | | | DKFZP434G032 | | | | | | | | , |
| rc AA024482 at | AA024482 | 45 | protein | #N/A | #N/A | 207.37 | 118.58 | 230.63 | 1. ن | -3.13 | 13.6 |
| rc_AA026030_at | AA026030 | 23 | EST | 11.01 | 0.01649 | 566.65 | 284.32 | 828.49 | 3.02 | 4.66 | 40.87 |
| r } | | | tyrosyl-tRNA | | | | | | | | |
| rc AA026270 at | AA026270 | 56 | synthetase | #N/A | #N/A | 201.1 | 224.51 | 50.82 | 76.85 | 20.92 | 45.22 |
| i I | | | olutamate-cysteine | | | | | | | | |
| | | | ligase (gamma- | | | | | | | | |
| | | | olutamyleyeteine | | | | | | | | |
| | | | glutal il ytoystell te evinthatasa) | | | | | | | | |
| | 7 | ò | symmetase), | V/1V# | V/14# | 70 00 | 24 37 | 13 20 | 12 50 | 12 27 | 8 96 |
| rc_AA035540_at | AAU35540 | ຂ່ | regulatory (50.6kiz) | ¥/N# | 4/N# | 59.04 | 10.40 | 13.23 | 0.00 | 12.21 | 200 |
| rc_AA040465_at | AA040465 | 92 | EST | 3.25 | 0.00146 | 161.91 | 125.35 | (2.19 | 47.13 | 40.00 | 0.40 |
| rc AA043944 at | AA043944 | 100 | EST | W/V# | #N/A | 52.9 | 29.13 | 46.43 | 3.69 | 3.15 | 7.26 |
| rc_AA053033_at | AA053033 | 124 | EST | 7.83 | 0.00379 | 212.61 | 135.28 | 160.82 | 3.41 | -1.28 | 19.74 |
| 1 | | | cadherin 17, Ll | | | | | | | | |
| rc 04053102 s at | AA053102 | 125 | intestine) | 26.63 | 0.01745 | 1053.43 | 972.71 | 785.51 | 4.92 | 4.94 | 9.04 |
| rc AA053248 i at | AA053248 | 128 | EST | 7.01 | 0.00003 | 2234.44 | 1595.15 | 1053.3 | 341.06 | 304.8 | 189.89 |
| r AA053660 at | AA053660 | 128 | EST | 15.98 | 0.00003 | 1573.49 | 1340 | 820.5 | 95.24 | 79.97 | 42.99 |
| | AAAAAAAA | 2 5 | - 53 | 42 83 | 0.00142 | 1270.03 | 1026 97 | 1179 12 | -1111 | -10.85 | 20.27 |
| rc_AAU338U3_s_at | AAUSSOUS | 132 | - 61 | 47.03 | 71.00.0 | 0.00 | 1050:01 | | | 2 | |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | fold | | | | | norma | norma | normal |
|------------------|----------|--------|---|----------|---------|-------------|-------------------------|-------------|--------|--------|---------|
| | | | | change | | metastatic: | metastatic: metastatic: | metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p.value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | glycoprotein A33 | | • | | | | | | |
| rc_AA055811_s_at | AA055811 | 133 | (transmembrane) collagen, type V, alpha | 6.86 | 0.02152 | 236.18 | 196.53 | 171.06 | 14.84 | 17.82 | 17.5 |
| rc AA055896 at | AA055896 | 135 | · · | 18.16 | 0.00146 | 540.67 | 538.43 | 382 | -3.14 | -5.8 | 33.88 |
| rc_AA084343_at | AA084343 | 178 | EST | #N/A | #N/A | 133.86 | 126.19 | 19.99 | 84.58 | 76.82 | 41.45 |
| ı | | | carcinoembryonic antigen-related cell | | | | | | | | |
| | | | adhesion molecule 6 | | | | ٠ | | | | |
| rc_AA100719_s_at | AA100719 | 212 | reacting antigen) | 50.88 | 0.00081 | 1405.2 | 1264.79 | 1137.97 | -17.03 | -22.23 | 16.43 |
| rc_AA112979_s_at | AA112979 | 225 | kinase 1 | #N/A | #N/A | 25.08 | 24.98 | 13.02 | 8.54 | 8.02 | 2.59 |
| rc AA115979 at | AA115979 | 232 | coil related protein | #N/A | #N/A | 74.58 | 89.38 | 66.52 | 22.91 | 13.67 | 29.42 |
| rc_AA126041_at | AA126041 | 244 | EST | #N/A | #N/A | 42.91 | 33.83 | 16.2 | 26.42 | 28.92 | 9.81 |
| AA127712_at | AA127712 | 255 | EST | 6.59 | 0.03706 | 332.45 | 208.85 | 296.68 | -29.54 | -28.71 | 104.37 |
| | | | amyloid beta (A4) precursor protein (protease nexin-II. | | | | | | | | |
| rc_AA128553 f_at | AA128553 | 260 | Alzheimer disease) | #N/A | #N/A | 87.35 | 102.82 | 24.17 | 114.91 | 108.16 | 30.28 |
| rc_AA131162_s_at | AA131162 | 266 | EST | 4.68 | 0.00042 | 117.43 | 123.78 | 39.05 | 11.86 | 13.57 | 19.83 |
| rc_AA134968_at | AA134968 | 289 | EST | 12.11 | 0.00079 | 322.34 | 208.47 | 256.66 | 10.04 | 7.96 | 15.53 |
| rc_AA150053_at | AA150053 | 327 | EST | 3.71 | 0.00102 | 309.61 | 304.16 | 82.48 | 101.87 | 88.37 | 73.73 |
| | | Ç | matrix | ļ | 0 | ļ | 3 | Š | ć | 9 | 9 |
| rc_AA151428_s_at | AA151428 | 333 | metalloproteinase 236 | cr.) | 0.0000 | 1/4.// | 131.00 | 104.51 | 3.23 | 10.01 | 24.39 |
| rc_AA156243_at | AA156243 | 340 | serine protease, umbilical endothelium | 41.44 | 0.00139 | 1255.4 | 547.15 | 1189.26 | 9.32 | 11.44 | 9.39 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | fold | | | | | normal | normal | normal |
|------------------|----------|--------|--------------------------|----------|---------|-------------|-------------------------|-------------|--------|--------|---------|
| | | | | change | _ | metastatic: | metastatic: metastatic: | metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | Mean | Medlan | Std Dev | Mean | Median | Std Dev |
| rc AA157857 s at | AA157857 | 350 | keratin 19 | #N/A | #N/A | 2728.74 | 2917.98 | 1076.95 | -13.21 | -14.4 | 15.84 |
| rc AA159525 at | AA159525 | 354 | EST | 49.39 | 0.00062 | 1309.05 | 1107.08 | 965.67 | -32.72 | -28.32 | 20.11 |
| rc_AA161043_at | AA161043 | 356 | tetraspan 1 | 18.8 | 0.00015 | 1477.19 | 932.02 | 1467.59 | 70.82 | 58.62 | 42.88 |
| rc_AA171760_at | AA171760 | 367 | EST | 17.86 | 0 | 645.43 | 578.38 | 312.37 | 28.03 | 16.24 | 57.21 |
| l I | | | polyglutamine binding | | | | | | | | |
| rc AA179787 at | AA179787 | 380 | protein 1 | 6.44 | 0.00206 | 191.33 | 171.71 | 138.9 | -11.56 | -12.46 | 42.46 |
| rc_AA179845_at | | 381 | EST | 5.77 | 0.01414 | 280.06 | 335.41 | 172.62 | 37.43 | 37.69 | 14.87 |
| rc_AA181600_at | | 384 | EST | 5.38 | 0.03316 | 166.88 | 94.16 | 153.49 | -40.51 | -47.81 | 24.34 |
| rc_AA196790_at | AA196790 | 421 | EST | 7.64 | 0.00287 | 239.45 | 275.18 | 142.37 | 22.91 | 26.8 | 10.54 |
| rc AA211483 at | AA211483 | 435 | EST | 44.07 | 0.00175 | 1300.23 | 1303.61 | 1051.58 | -29.25 | -34.99 | 17.49 |
| rc_AA227926_at | AA227926 | 460 | EST | 6.81 | 0.01701 | 228.91 | 120.1 | 243.92 | 16.24 | 14.21 | 98.9 |
| rc_AA232508_at | AA232508 | 464 | EST | #N/A | #N/A | 464 | 533.98 | 266.87 | 170.48 | 156.89 | 104.75 |
| l | | | general transcription | | | | | | | | |
| | | | factor IIIC, polypeptide | | | | | | | | |
| rc AA233290 at | AA233290 | 469 | 220kD) | #N/A | #N/A | 56.54 | 43.91 | 22.15 | 28.32 | 31.59 | 25.36 |
| rc_AA234096_at | AA234096 | 479 | EST | 23.72 | 0.00018 | 607.06 | 420.34 | 366.53 | -3.47 | 1.73 | 30.91 |
| rc_AA234346_at | AA234346 | 480 | EST | #N/A | #N/A | 36.98 | 39.47 | 12.81 | 6.21 | 6.79 | 19.99 |
| rc_AA234362_at | AA234362 | 481 | EST | 3.89 | 0.03524 | 116.26 | 75.37 | 105.77 | -1.86 | -4.2 | 16.67 |
| | | | survival of motor | | | | | | | | |
| | | | neuron protein | | | | | | | | |
| rc_AA234365_at | AA234365 | 482 | interacting protein 1 | #N/A | #N/A | 56.37 | 42.6 | 34.73 | 12.96 | 16.93 | 12.99 |
| rc_AA234706_at | AA234706 | 488 | EST | #N/A | W/A# | 68.33 | 53.21 | 21.55 | 49.66 | 39.76 | 42.06 |
| rc_AA235707_at | AA235707 | 200 | EST | 9.17 | 0.00005 | 189.42 | 161.88 | 57.05 | -18.19 | -19.06 | 14.8 |
| 1 | | | ecotropic viral | | | | | | | | |
| rc AA236533 s at | AA236533 | 514 | integration site 1 | 4.01 | 0.02882 | 106.15 | 76.05 | 78.18 | -16.45 | -14.04 | 8.52 |
| rc_AA243173_at | AA243173 | 526 | EST | 8.75 | 0.00003 | 384.86 | 482.53 | 162.08 | 41.66 | 40.74 | 27.31 |
| | | | apoptosis inhibitor 4 | | | | | | | | |
| rc_AA252994_at | AA252994 | 222 | (survivin) | 3.55 | 0.00075 | 152.63 | 140.53 | 31.01 | 50.23 | 40.59 | 35.61 |
| | | | | | | | | | | | |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | fold | | | | | normal | normal | normal |
|------------------|----------|--------|-------------------------------------|----------|---------|-------------|-------------------------|-------------|--------|---------|------------------|
| | | | | change | | metastatic: | metastatic: metastatlc: | metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Mrown Gene Name | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | adaptor-related | | | | - | | | | : : : : |
| AA253330 s at | AA253330 | 562 | gamma 1 subunit | 3.87 | 0.00708 | 219.95 | 186.12 | 141.92 | 47.32 | 51.59 | 30.28 |
| rc AA253473 at | AA253473 | 267 | EST | 15.23 | 0.00171 | 375.2 | 435.68 | 216.18 | -4.38 | -3.19 | 12.35 |
| | AA255566 | 570 | EST | #N/A | #N/A | 63.43 | 65.2 | 30.17 | 16.18 | 14.38 | 13.17 |
| rc_AA256642_at | AA256642 | 582 | EST | 11.17 | 0.00035 | 266.31 | 275.43 | 135.85 | 0.78 | -1.34 | 15.74 |
| i | | | WW domain binding protein 4 (formin | ū | | | | | | | |
| rc AA258836 at | AA258836 | 601 | binding protein 21) | #N/A | #N/A | 41.9 | 44.58 | 13.6 | 26.08 | 23.64 | 10.4 |
| rc_AA262943_at | AA262943 | 611 | EST | 13.42 | 0.00234 | 430.7 | 207.29 | 511.01 | -1.08 | -6.93 | 19.06 |
| | | | neuroblastoma- | | | | | | | | |
| rc_AA281214_s_at | AA281214 | 643 | amplified protein | #N/A | #N/A | 130.98 | 133.63 | 68.42 | 68.69 | 53.91 | 39.31 |
| rc_AA282505_at | AA282505 | 629 | EST | #N/A | #N/A | 461.54 | 478.83 | 622.39 | -41.45 | -17 | 98.17 |
| | | | putative G protein- | | | | | | | | |
| rc_AA291259_at | AA291259 | 697 | coupled receptor | #N/A | #N/A | 112.28 | 45.1 | 113.44 | 1.53 | 1.88 | 90.9 |
| AA291456_s_at | AA291456 | 200 | EST | 3.96 | 0.03633 | 1808.4 | 1273.75 | 1481.84 | 671.21 | 509.58 | 629.13 |
| rc_AA292765_at | AA292765 | 712 | ZW10 interactor | 7.14 | 0.02623 | 230.83 | 166.72 | 232.16 | 11.33 | 18.29 | 13.27 |
| rc_AA292931_at | AA292931 | 715 | EST | 3.97 | 0.00067 | 230.65 | 260.43 | 79.42 | 57.32 | 61.89 | 18.62 |
| AA295819_s_at | AA295819 | 722 | EST | ထ | 0.01793 | 230.14 | 251.43 | 146.89 | -14.93 | -20.8 | 19.72 |
| rc_AA331393_at | AA331393 | 739 | EST | 16.73 | 0.00848 | 600.02 | 315.98 | 598.52 | 5.08 | 3.88 | 9.43 |
| rc_AA335191_f_at | AA335191 | 741 | creatine kinase, brain | 47.35 | 0.00419 | 3540.65 | 3971.43 | 3135.19 | 43.84 | 33.21 | 43.52 |
| rc AA349792 s at | AA349792 | 759 | homolog | W/W# | #N/A | 525.17 | 553.97 | 245.2 | 221.5 | 212.24 | 98.86 |
| rc_AA372018_at | AA372018 | 768 | EST | 14.3 | 0.00178 | 363.44 | 220.03 | 261.04 | -2.44 | -3.09 | 8.36 |
| ı | | | differentially expressed in | | | | | | | | |
| AA372630 s at | AA372630 | 769 | lineanes | 25 49 | 0.01743 | 2004.32 | 436 62 | 3873 61 | 6.67 | 4.87 | 7 16 |
| in | 22041200 | 3 | 20000 | 7.77 | 2.00 | 4004.04 | 40.004 | 20.0 | 5 | į. Ž | 2 |
| | | | | | | | | | | | |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | fold | | | | | normal | normal | normal |
|----------------|------------|--------|--------------------------------------|----------|---------|-------------|-------------------------|-------------|---------|---------|---------|
| | | | | change | | metastatic: | metastatic: metastatic: | metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | DKFZP586B0519 | | | | | | | | |
| AA384184_s_at | AA384184 | 774 | protein | 3.38 | 0.01209 | 476.17 | 411.64 | 148.86 | 181.02 | 188.54 | 135.4 |
|]] | | | laminin receptor 1 | | | | | | | | |
| | | | (67kD, ribosomal | | | | | | | | |
| rc_AA394121_at | AA394121 | 778 | protein SA) | 23.78 | 0.00099 | 788.51 | 677.33 | 612.74 | 0.21 | -8.38 | 31.59 |
| rc_AA398908_at | AA398908 | 801 | EST | 38.69 | 0.01089 | 1678.58 | 1796.68 | 1500.48 | -174.29 | -216.08 | 104.42 |
| l | | | tight junction protein 3 | | | | | | | | |
| rc_AA399226_at | AA399226 | 803 | (zona occludens 3) | 3.59 | 0.02002 | 89.54 | 78.97 | 63.98 | -5.01 | -6.48 | 11.33 |
| l | | | tumor suppressor | | | | | | - | | |
| | | | deleted in oral cancer- | | | | | | | | |
| rc AA401965 at | AA401965 | 833 | related 1 | #N/A | #N/A | 384.12 | 271.95 | 444.06 | 120.58 | 93.29 | 109.41 |
| rc AA404597 at | AA404597 | 854 | EST | #N/A | #N/A | 624.37 | 495.56 | 274.8 | 379.26 | 336.33 | 167.43 |
| rc_AA405715_at | AA405715 | 862 | hypothetical protein | 4.68 | 0.00898 | 152.23 | 138.92 | 96.78 | 24.47 | 27.3 | 11.75 |
| rc AA406542 at | AA406542 | 878 | EST | 8.27 | 0.00724 | 230.18 | 170.94 | 206.8 | -1.99 | 1.49 | 12.75 |
| rc_AA410469_at | AA410469 | 883 | EST | 6.3 | 0.00103 | 337.03 | 250.11 | 264.1 | 50.51 | 42.79 | 60.18 |
| i I | | | DKFZP586E1422 | | | | | | | | |
| rc_AA419217_at | AA419217 | 923 | protein | 6.77 | 0.00045 | 276.53 | 215.37 | 172.25 | 36.93 | 36.7 | 21.88 |
| l I | | | anterior gradient 2 (Xenepus laevis) | | | | | | | | |
| rc AA421562 at | - AA421562 | 934 | homolog | 56.3 | 0.0041 | 2556.78 | 792 | 3323.39 | 14.22 | 15.72 | 6.54 |
| rc_AA422086_at | AA422086 | 938 | EST | 10.71 | 0.03418 | 828.27 | 598.22 | 800.79 | 39.48 | 34.98 | 23.29 |
| l I | | | cytochrome P540 family member | | | | | | | | |
| rc AA422150 at | AA422150 | 939 | predicted from ESTs | 17.14 | 0.00108 | 664.26 | 606.609 | 475.91 | 28.76 | 27.53 | 22.47 |
| rc_AA424487_at | AA424487 | 945 | EST | 38.41 | 0.00002 | 2689.32 | 1863.02 | 1900.51 | 67.92 | 61.95 | 46.24 |
| rc AA424881 at | AA424881 | 949 | EST | 6.3 | 0.00556 | 158.13 | 155.45 | 82.63 | -16.85 | -22.15 | 20.49 |
| rc_AA425279_at | AA425279 | 951 | quiescin Q6 | 6.15 | 0.00083 | 221.93 | 181.8 | 167.44 | 18.33 | 22.33 | 44.81 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | fold | | | | | normal | normal | normal |
|--|-----------------|--------|---|----------|---------|-------------|-------------------------|-------------|--------|---------|----------------|
| | | | | change | | metastatic: | metastatic: metastatic: | metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | serine/threonine kinase 24 (Ste20, | , | | | | | | | |
| rc_AA425401_at | AA425401 | 954 | yeast homolog) | 3.22 | 0.00625 | 246.6 | 166.4 | 175.99 | 76.4 | 78.33 | 43.72 |
| rc_AA425852_i_at | AA425852 | 928 | EST | 8. 8. | 0.03874 | 149.8 | 125.7 | 128.03 | 7.02 | 5.22 | 4.75 |
| | | | ATP-binding cassette, sub-family F (GCN20). | | | | | | | | |
| rc_AA427460_s_at | AA427460 | 972 | member 2 | #N/A | #N/A | 215.08 | 135.22 | 216.95 | 92.5 | 83.63 | 49.3 |
| AA427468_s_at | AA427468 | 973 | claudin 4 | 84.43 | 0 | 5646.4 | 5344.98 | 1581.19 | 72.58 | 64.85 | 87.74 |
| rc_AA427636_at | AA427636 | 976 | EST | 19.23 | 0.00145 | 511.98 | 500.83 | 299.69 | 6.19 | 3.7 | 15.84 |
| , | | | serine protease | | | | | | ٠ | | |
| rc_AA429009_at | AA429009 | 994 | inhibitor, Kunitz type 1 | 30.04 | 0.00001 | 1010.32 | 1100.19 | 472.23 | -2.97 | -24.67 | 62.21 |
| te a 0080000 21 | 00800 | 1001 | displatin resistance associated | 12 51 | 0.00053 | 1925 79 | 500.00 | 77 600 | 60 00 | 67 23 | 50 25 |
| 10_7423030_3_at | A A 420674 | 2 5 | associated FOT | 2.7 | | 454.02 | 306.45 | 370 26 | 20.02 | 00.00 | 59.55 |
| 10_A44500/4_at | AA430074 | 1056 | EST. | 2.5. | 0.00293 | 70.05 | 506.15 04.2E | 57 6.30 | 5.7. | 2 24 | 39.30 4F.26 |
| 10 A443300 10 at | 4440010 | | [5] | 5 5 | 0.04402 | 19:30 | 91.33 | 02.03 | 2 3 | 47.0 | 10.30 |
| rc_AA442763_ar | AA442/63 | 701 | cyclin bz | 5.03 | 0.02168 | 136.16 | 109.91 | 91.52 | -14.64 | -15.1 | 13.08 |
| | | | sarroma viral | | | | ,. | | | | |
| rc AA443316 s at | AA443316 | 1075 | oncogene homolog | 4.13 | 0.01729 | 191.06 | 173.3 | 113.42 | 40.07 | 41.12 | 32.54 |
| 1 1 | | | tumor suppressing | | | | | | | | |
| rc AA443941 at | AA443941 | 1085 | candidate 1 | 3.57 | 0.01685 | 137.02 | 138 22 | 83.35 | 31.00 | 34.0 | 16.61 |
| rc AA447687 at | AA447687 | 1104 | FST | 11 42 | 0.00362 | 30. 908 | 178.65 | 244 12 | -16 27 | -14 45 | 8 55 |
| rc AA447991 at | A A 4 4 7 9 9 1 | 1112 | 183 | 4 00 | 0.00173 | 279 70 | 234 63 | 161 24 | 64 40 | 40.73 | 53 34 |
| rc AAAAAKE at | AAAAAAE | 1106 | FOT | 200 | 0.000 | 01.01.2 | 775 84 | 377 90 | 160 02 | 178 76 | 113 74 |
| 0 000000000000000000000000000000000000 | AA64077 | 1120 | FST | 0.43 | 0.0000 | 73000 | 157.2 | 66.776 | 27.20 | 27.50 | 10.70 |
| 1C_A4451077_at | - | 001 | - C - C - C - C - C - C - C - C - C - C | 0.00 | 0.00409 | 239.33 | 5.75 | 244.02 | 07:17- | EC. 12- | 10.20 |
| rc_AA454597_s_at | AA454597 | 1166 | ESI | 3.63 | 0.0067 | 268.15 | 274.94 | 167.72 | 9.99 | 57.29 | 23.41 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| Affy ID Genbank Seq ID Known Gene Name in metas p value metastatic: metastati | | | | | fold | | | | | normal | normal | normal |
|---|------------------|----------|--------|-----------------------|----------|---------|-------------|--------------|-------------|--------|--------|---------|
| Genbank Seq ID Known Gene Name in metas p value Mean Median S AA454908 171 product 9.3 0.06539 320.03 253.32 263.32 4A455226 120.0 EST 4MINA #MINA 481.26 138.32 4A455226 120.0 EST 4MINA #MINA 461.02 58.13 4A456366 123.12 EST 4MINA #MINA 461.02 58.13 4A466366 127.22 311.71 6 4A466362 120.1 EST 4MINA #MINA #MINA 4MINA | | | | | change | | metastatic: | | metastatic: | set 2: | set 2: | set 2: |
| KiAA0144 gene | Affy ID | Genbank | Seq ID | 줃 | in metas | | Mean | Median | Std Dev | Mean | Median | Std Dev |
| AA45908 1171 product 9.3 0.00539 320.03 253.32 AA457235 1200 EST #NIA #NIA 481.26 138.32 48.485 AA45924 121 EST #NIA #NIA 481.26 138.32 48.485 AA46962 1251 EST #NIA #NIA 61.02 311.71 64.485 AA465361 1251 EST #NIA #NIA 61.02 56.49 58.13 AA465360 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA46560 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA46560 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA46560 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA46560 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA46560 128 me | | | | | | | | | | | | |
| AA457235 1200 EST #N/A #N/A #R/A 481.26 138.32 48.45 AA459254 1211 EST 5.36 0.0259 403.69 194.85 4.85 AA469666 1231 EST #N/A #N/A #N/A 56.19 58.13 AA463861 1251 EST #N/A #N/A 41.02 56.4 AA465690 1271 EST #N/A #N/A 171.26 118.15 Lumor necrosis factor receptor superfamily, #N/A #N/A 171.26 118.15 AA465690 1273 LIM domain binding 1 #N/A #N/A 171.26 118.15 AA45660 1271 EST #N/A #N/A 171.26 118.15 AA476649 1286 member 12 #N/A #N/A 172.35 44.04 AA477561 1292 EST #N/A #N/A 172.35 14.04 AA477561 1292 EST #N/A #N/A | AA454908 s at | AA454908 | 1171 | product | 9.3 | 0.00539 | 320.03 | 253.32 | 238.5 | -33.7 | -46.73 | 73.56 |
| AA459254 1211 EST 5.36 0.0259 403.69 194.85 AA460666 1231 EST #NIA #NIA 56.19 58.13 AA463861 1251 EST #NIA #NIA 61.02 56.4 AA465342 1271 EST #NIA #NIA 61.02 56.4 AA465660 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA465660 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA465660 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA465660 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA4776749 1286 membrane protein) #NIA #NIA 160.87 104.95 AA477561 1292 EST #NIA #NIA 172.36 140.4 AA482546 1336 KIAA0124 protein #NIA #NIA 143.86 | rc AA457235 at | AA457235 | 1200 | EST | #N/A | #N/A | 481.26 | 138.32 | 573.73 | 11.06 | 17.32 | 15.26 |
| AA46566 1231 EST #NIA #NIA 56.19 58.13 AA463861 1251 EST #NIA #NIA 61.02 56.4 AA46560 1273 LIM domain binding 1 #NIA #NIA 61.02 56.4 AA46560 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA46560 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA46560 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA46560 1273 LIM domain binding 1 #NIA #NIA 171.26 118.15 AA476749 1286 membrane protein) #NIA #NIA 160.87 104.95 AA477561 1292 EST #NIA #NIA 160.87 104.95 AA477561 1292 EST #NIA #NIA 152.35 136.55 AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA491208 1388 reading frame 1 #NIA #NIA 159.88 143.86 AA59958 1454 EST #NIA #NIA 143.86 AA59958 1454 EST #NIA #NIA 143.86 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA60905 1502 EST #NIA #NIA 147.72 165.83 AA62095 1512 EST #NIA #NIA 147.72 165.83 AA62095 1512 EST #NIA #NIA 147.72 165.83 AA62095 1512 EST #NIA #NIA 147.72 165.83 AA6202304 1530 EST #NIA #NIA 147.72 165.83 AA621277 1520 EST #NIA #NIA 160.71 160.71 80.15 AA621277 1520 EST #NIA #NIA 584.5 621.94 | rc_AA459254_at | AA459254 | 1211 | EST | 5.36 | 0.0259 | 403.69 | 194.85 | 401.21 | 51.84 | 43.51 | 37.57 |
| AA463861 1251 EST 24.79 0.00096 672.22 311.71 644 AA465342 1271 EST #NI/A #NI/A #NI/A 113.36 100.15 AA465660 1273 LIM domain binding 1 #NI/A #NI/A 171.26 118.15 AA465660 1273 LIM domain binding 1 #NI/A #NI/A 171.26 118.15 Immor necrosis factor receptor superfamily, receptor superfamily, 171.26 118.15 member 12 (translocating chain-association) #NI/A #NI/A 160.87 104.95 AA477561 1292 EST #NI/A #NI/A 172.35 136.55 AA482546 1336 IRA0124 protein 4.41 0.00604 181.72 113.91 AA4812546 1336 IRA0124 protein #NI/A #NI/A #NI/A 77.48 78.85 AA599244 1448 KIAA0530 protein #NI/A #NI/A #NI/A #NI/A #NI/A #NI/A #NI/A #NI/ | rc_AA460666_at | AA460666 | 1231 | EST | #N/A | #N/A | 56.19 | 58.13 | 26.29 | -24.55 | -17.23 | 20.98 |
| AA464962 1264 EST #NI/A #NI/A #10.2 56.4 AA465342 1271 EST #NI/A #NI/A 113.36 100.15 AA465660 1273 LIM domain binding 1 #NI/A #NI/A 171.26 118.15 Immor necrosis factor receptor superfamily, receptor superfamily, 171.26 118.15 member 12 (translocating chain-association) association 44.04 44.04 AA477561 1292 EST #NI/A #NI/A 172.35 136.55 AA477561 1292 EST #NI/A #NI/A 172.35 136.55 AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA4892546 1336 KIAA0530 protein 4.41 0.00604 181.72 143.86 AA599244 1448 KIAA0530 protein 4.41 0.001246 77.48 78.95 AA609913 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 | rc_AA463861_at | AA463861 | 1251 | EST | 24.79 | 0.00096 | 672.22 | 311.71 | 641.54 | -22.02 | -20.04 | 16.5 |
| AA465342 1271 EST #NI/A #NI/A #11.36 100.15 AA465660 1273 LIM domain binding 1 #NI/A #NI/A 171.26 118.15 receptor superfamily, member 12 receptor superfamily, member 12 (translocating chain-association association | rc_AA464962_at | AA464962 | 1264 | EST | #N/A | #N/A | 61.02 | 56.4 | 30.98 | 20.99 | 14.37 | 19.64 |
| AA465660 1273 LIM domain binding 1 m/l/A m/l/A m/l/A m/l/A momber 12 (translocating chain-association association member 12 (translocating chain-association member 12 (translocating chain-association association membrane protein) membrane protein) membrane protein association association membrane protein association membrane protein association membrane protein membrane protein membrane protein membrane protein membrane protein association association association association association membrane protein association associatio | rc_AA465342_at | AA465342 | 1271 | EST | #N/A | #N/A | 113.36 | 100.15 | 87.62 | 30.07 | 33.06 | 23.97 |
| tumor necrosis factor receptor superfamily, member 12 (translocating chain-association AA476749 1286 membrane protein) #N/A #N/A 160.87 104.95 AA478556 1303 EST AA482546 1336 KIAA0124 protein AA482546 1338 KIAA0124 protein AA491208 1388 reading frame 1 AA491208 1388 reading frame 1 #N/A #N/A 159.88 143.86 AA599244 1448 KIAA0530 protein #N/A #N/A 159.88 143.86 AA609013 1477 dipeptidase 1 (renal) #N/A #N/A 66.17 48.91 AA609095 1512 EST #N/A #N/A 147.72 165.83 AA620497 1503 EST #N/A #N/A 147.72 165.83 AA620497 1520 EST #N/A #N/A 147.72 165.83 AA620497 1520 EST #N/A #N/A 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #N/A #N/A 584.5 621.94 | rc_AA465660_at | AA465660 | 1273 | LIM domain binding 1 | #N/A | #N/A | 171.26 | 118.15 | 103.39 | 124.59 | 105.93 | 56.71 |
| receptor superfamily, member 12 (translocating chain-association (translocating chain-association) #N/A #N/A 53.23 44.04 AA477561 1286 membrane protein) #N/A #N/A 104.95 AA478556 1303 EST 44.1 0.00604 181.72 104.95 AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA482546 1336 KIAA0124 protein #N/A #N/A 181.72 113.91 AA482546 1336 KIAA0530 protein #N/A #N/A 159.88 143.86 AA599585 1454 EST #N/A #N/A 77.48 78.85 AA699585 1454 EST #N/A #N/A 147.72 165.83 AA609013 1477 EST #N/A #N/A 147.72 165.83 AA620995 1512 EST | l I | | | tumor necrosis factor | | | | | | | | |
| (translocating chain-association #N/A #N/A #S3.23 44.04 AA476749 1286 membrane protein) #N/A #N/A 160.87 104.95 AA47856 1303 EST #N/A #N/A 160.87 104.95 AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA491208 138 reading frame 1 #N/A #N/A 159.88 143.86 AA599244 1448 KIAA0530 protein #N/A #N/A 66.17 48.91 AA599585 1454 EST #N/A #N/A 147.72 165.83 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA620497 1503 EST #N/A #N/A #N/A 100.71 80.15 AA620956 1512 EST 3.74 0.03414 100.71 80.15 AA621277 1520 EST 3.81 0.00194 <td></td> <td></td> <td></td> <td>receptor superfamily,</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> | | | | receptor superfamily, | | | | | | | | |
| AA476749 1286 membrane protein) #N/A #N/A 53.23 44.04 AA476549 1286 membrane protein) #N/A #N/A 160.87 104.95 AA47856 1303 EST #N/A #N/A 172.35 136.55 AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA491208 1388 reading frame 1 #N/A #N/A 159.88 143.86 AA599244 1448 KIAA0530 protein 3.39 0.01246 77.48 78.85 AA599585 1454 EST #N/A #N/A 66.17 48.91 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA620497 1503 EST #N/A #N/A 147.72 165.83 AA620995 1512 EST #N/A 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 <td></td> <td></td> <td></td> <td>member 12</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> | | | | member 12 | | | | | | | | |
| AA476749 1286 membrane protein) #N/A #N/A 53.23 44.04 AA477561 1292 EST #N/A #N/A 160.87 104.95 AA482546 1303 EST #N/A #N/A 172.35 136.55 AA482546 1303 EST #N/A #N/A 172.35 136.55 AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA491208 1388 reading frame 1 #N/A #N/A 159.88 143.86 AA599585 1454 EST #N/A #N/A 66.17 48.91 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA620497 1503 EST #N/A #N/A 147.72 165.83 AA620497 150 EST #N/A 100.71 80.15 AA621277 1520 EST 3.74 0.003414 100.71 80.15 AA | | | | (translocating chain- | | | | | | | | |
| AA476749 1286 membrane protein) #N/A #N/A 53.23 44.04 AA477561 1292 EST #N/A #N/A 160.87 104.95 AA478556 1303 EST #N/A #N/A 172.35 136.55 AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA491208 1388 reading frame 1 #N/A #N/A 159.88 143.86 AA599585 1454 EST #N/A #N/A 66.17 48.91 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA609795 1492 EST #N/A #N/A 147.72 165.83 AA620497 1503 EST #N/A 100.71 80.15 AA621277 1520 EST 3.74 0.00194 81.18 74.81 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB | | | | association | | | | | | | | |
| AA477561 1292 EST #NIA #NIA #NIA 160.87 104.95 AA478556 1303 EST #NIA #NIA 172.35 136.55 AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA491208 1388 reading frame 1 #NIA #NIA 159.88 143.86 AA599585 1454 EST #NIA #NIA 66.17 48.91 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA609795 1492 EST #NIA #NIA 147.72 165.83 AA620497 1503 EST #NIA 100.71 80.15 AA621277 1520 EST 3.74 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #NIA #NIA 584.5 621.94 | rc AA476749 s at | AA476749 | 1286 | membrane protein) | #N/A | #N/A | 53.23 | 44.04 | 18.16 | 25.5 | 29.62 | 11.96 |
| AA478556 1303 EST #NIA #NIA 172.35 136.55 AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA491208 1388 reading frame 1 #NIA #NIA 159.88 143.86 AA599585 1454 EST #NIA #NIA 48.91 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA609795 1492 EST #NIA #NIA 147.72 165.83 AA620497 1503 EST #NIA #NIA 100.71 80.15 AA621277 1520 EST 3.74 0.03414 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #NIA #NIA 584.5 621.94 | rc AA477561 at | AA477561 | 1292 | EST | ∀/Ń# | #N/A | 160.87 | 104.95 | 133.7 | 63.83 | 76.56 | 37.87 |
| AA482546 1336 KIAA0124 protein 4.41 0.00604 181.72 113.91 AA491208 1388 reading frame 1 #N/A #N/A 159.88 143.86 AA599244 1448 KIAA0530 protein 3.39 0.01246 77.48 78.85 AA599585 1454 EST #N/A #N/A 66.17 48.91 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA620497 1503 EST #N/A #N/A 147.72 165.83 AA620497 1502 EST 3.74 0.03414 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #N/A #N/A 584.5 621.94 | rc_AA478556_at | AA478556 | 1303 | EST | #N/A | #N/A | 172.35 | 136.55 | 120.49 | 143.25 | 134.62 | 109.1 |
| AA491208 1388 reading frame 1 #N/A #N/A #159.88 143.86 AA599244 1448 KIAA0530 protein 3.39 0.01246 77.48 78.85 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA609795 1492 EST #N/A #N/A 74.38 34 AA620497 1503 EST #N/A #N/A 147.72 165.83 AA621277 1520 EST 3.74 0.03414 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #N/A #N/A 584.5 621.94 | rc AA482546 s at | AA482546 | 1336 | KIAA0124 protein | 4.41 | 0.00604 | 181.72 | 113.91 | 140.73 | 38.26 | 30.26 | 32.72 |
| AA491208 1388 reading frame 1 #NI/A #NI/A 159.88 143.86 AA599244 1448 KIAA0530 protein 3.39 0.01246 77.48 78.85 AA599585 1454 EST #NI/A #NI/A 66.17 48.91 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA609795 1492 EST #NI/A #NI/A 74.38 34 AA620497 1503 EST #NI/A #NI/A 147.72 165.83 AA621277 1520 EST 3.74 0.03414 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #NI/A #NI/A 584.5 621.94 | ! ! | | | chromosome 6 open | | | | | | | | |
| AA599244 1448 KIAA0530 protein 3.39 0.01246 77.48 78.85 AA599585 1454 EST #NIA #NIA 66.17 48.91 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA609795 1492 EST #NIA #NIA 147.72 165.83 AA620497 1503 EST 3.74 0.03414 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #NIA #NIA 584.5 621.94 | rc AA491208 at | AA491208 | 1388 | _ | W/V# | #N/A | 159.88 | 143.86 | 80.49 | 123.37 | 131.81 | 47.97 |
| AA599585 1454 EST #NI/A #NI/A 66.17 48.91 AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA609795 1492 EST #NI/A #NI/A 74.38 34 AA620497 1503 EST #NI/A #NI/A 147.72 165.83 AA621277 1520 EST 3.74 0.03414 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #NI/A #NI/A 584.5 621.94 | rc AA599244 at | AA599244 | 1448 | _ | 3.39 | 0.01246 | 77.48 | 78.85 | 37.98 | 7.71 | 90.9 | 7.11 |
| AA609013 1477 dipeptidase 1 (renal) 10.17 0.00109 1185.55 1222.89 AA609795 1492 EST #N/A #N/A 74.38 34 AA620497 1503 EST #N/A 147.72 165.83 AA620995 1512 EST 3.74 0.03414 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #N/A #N/A 584.5 621.94 | rc_AA599585_at | AA599585 | 1454 | | #N/A | #N/A | 66.17 | 48.91 | 48.12 | -12.47 | -17.55 | 21.82 |
| AA609795 1492 EST #N/A #N/A 74.38 34 AA620497 1503 EST #N/A #N/A 147.72 165.83 AA620995 1512 EST 3.74 0.03414 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #N/A #N/A 584.5 621.94 | rc AA609013 s at | AA609013 | 1477 | dipeptidase 1 (renal) | 10.17 | 0.00109 | 1185.55 | 1222.89 | 634.13 | 114.31 | 98.91 | 75.22 |
| AA620497 1503 EST #N/A #N/A 147.72 165.83 AA620995 1512 EST 3.74 0.03414 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #N/A #N/A 584.5 621.94 | rc AA609795 at | AA609795 | 1492 | | #N/A | #N/A | 74.38 | 8 | 77.76 | -41.44 | -45.5 | 28.82 |
| AA620995 1512 EST 3.74 0.03414 100.71 80.15 AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #N/A #N/A 584.5 621.94 | rc AA620497 s at | AA620497 | 1503 | | #N/A | #N/A | 147.72 | 165.83 | 66.02 | 95.91 | 76.7 | 80.33 |
| AA621277 1520 EST 3.81 0.00194 81.18 74.81 AB002304 1534 KIAA0306 protein #N/A #N/A 584.5 621.94 | rc AA620995 at | AA620995 | 1512 | | 3.74 | 0.03414 | 100.71 | 80.15 | 101.95 | 3.5 | 1.8 | 3.77 |
| t AB002304 1534 KIAA0306 protein #N/A #N/A 584.5 621.94 | rc_AA621277_at | AA621277 | 1520 | | 3.81 | 0.00194 | 81.18 | 74.81 | 30.84 | -0.15 | 1.98 | 12.84 |
| | AB002304_at | AB002304 | 1534 | ΧĀ | #N/A | #N/A | 584.5 | 621.94 | 136.64 | 313.99 | 298.88 | 88.6 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | fold | | - | | | normal | normal | normal |
|----------------|----------|--------|--------------------------------|----------|---------|-------------|-------------------------------------|-------------|--------|--------|---------|
| | | | - | change | _ | metastatic: | metastatic: metastatic: metastatic: | metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Sed ID | Genbank Seq ID Known Gene Name | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | KIAA0351 gene | | | | | | | | |
| AB002349_at | AB002349 | 1537 | product | W/N# | #N/A | 65.63 | 52.11 | 32.26 | 49.37 | 46.52 | 37.93 |
| | | | lectin, galactoside- | | | | | | • | , | |
| | | | binding, soluble, 9 | | | | | | | | |
| AB006782_at | AB006782 | 1541 | (galectin 9) | WWW. | #N/A | 475.47 | 430.44 | 192.68 | 222.25 | 213.37 | 51.25 |
| AF003521_at | AF003521 | 1545 | jagged 2 | 11.26 | 0.00008 | 426 | 330.95 | 309.91 | 15.48 | -3.6 | 91.81 |
| C00808_s_at | C00808 | 1553 | EST | #N/A | W/V# | 122.57 | 118.32 | 53.36 | 91.65 | 87.87 | 32.51 |
| C14228 f at | C14228 | 1567 | EST | Y/V# | #N/A | 39.93 | 33.15 | 13.59 | -0.51 | 2.85 | 8.61 |
| rc_C15078_i_at | C15078 | 1573 | EST | #N/A | #N/A | 115.08 | 73.82 | 91.34 | 26.22 | 31.14 | 60.2 |
| , | , | | pituitary tumor- | | | | | | | | |
| rc_C21248_at | C21248 | 1585 | transforming 1 | 3.85 | 0.00456 | 100.24 | 96.81 | 47.17 | -1.7 | -11.27 | 28.49 |
| | | | phosphatidylinositol | | | | | | | | , |
| D13435_at | D13435 | 1605 | glycan, class F | #N/A | #N/A | 80.8 | 88.58 | 41.56 | 34.85 | 41.76 | 18.11 |
| | | | basic transcription | | | | | | | | |
| | | | element binding | | | | | | | | |
| D14520_at | D14520 | 1613 | protein 2 | 4.93 | 0.00004 | 151.12 | 135.19 | 57.24 | 25.67 | 17.28 | 25.67 |
| rc_D20906_at | D20906 | 1627 | EST | 5.18 | 0.02189 | 210.11 | 151.67 | 185.24 | 14.87 | 11.71 | 27.2 |
| | | | minichromosome | | | | | | • | | |
| | | | maintenance deficient | | | | | | | | ÷ |
| | | | (S. cerevisiae) 2 | | | • | | | | | |
| D21063_at | D21063 | 1628 | (mitotin) | 3.83 | 0.00983 | 91.18 | 55.12 | 60.89 | -51,33 | -62.63 | 28.4 |
| D50914_at | D50914 | 1673 | KIAA0124 protein | 4.74 | 0.00752 | 116.55 | 91.64 | 89.27 | -17.4 | -17.86 | 16.97 |
| | | | KIAA0152 gene | | | | | | - | | |
| D63486_at | D63486 | 1712 | product | 6.33 | 0.00078 | 543.91 | 676.55 | 287.16 | 84.6 | 95.61 | 45.48 |
| | | | KIAA0159 gene | | | | | | | | |
| D63880_at | D63880 | 1715 | product | 4.26 | 0.00253 | 93.03 | 103.26 | 41.4 | -2.12 | 4.45 | 9.38 |
| D79992_at | D79992 | 1724 | product | #N/A | #N/A | 35.23 | 42.16 | 14.88 | 22 | 18.09 | 11.9 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| 1 | | | | | | | | | | | | | | | | | | | • | |
|---------------|---|---|--|---|--|------------------|--|--------------------|-------------------------|--|--|--|----------------------|--|--|--|---|---|--|--|
| | 13.07 | 24.59 | 17.46 | 57.58 | | 27.84 | | 28.15 | 21.13 | 17.79 | 58.6 | 47.16 | i | 9.79 | , | 13.52 | 35.81 | | 13.83 | |
| | 6.15 | 28.29 | 43.53 | 43.06 | | 19.65 | | 43.12 | 33.07 | 18.99 | 195.54 | 232.6 | ! | 17.26 | | 20.65 | -50.25 | | 20.29 | |
| | 8.57 | 22.95 | 41.25 | 42.54 | | 23.23 | | 37.22 | 33.34 | 23.85 | 178.42 | 235.37 | : | 14.92 | | 16.22 | -49.72 | | 19.92 | |
| | 53.87 | 149.08 | 42.83 | 172.25 | | 216.64 | | 102.81 | 13.36 | 47.7 | 83,95 | 142.63 | | 49.41 | | 262.03 | 631.83 | | 27.67 | |
| | 42.49 | 303.7 | 67.39 | 436.55 | | 156.39 | | 125.75 | 34.37 | 129.16 | 170.47 | 598.46 | | 115.96 | | 366.72 | 696.47 | | 112.31 | |
| Meall | 55.25 | 253.6 | 82.4 | 486.85 | | 236.64 | | 142.92 | 40 84 | 117.8 | 199.08 | 638.66 | | 90.37 | | 311.05 | 865.39 | | 104.92 | |
| n value | #N/A | 0.00213 | #N/A | 0.00001 | | 0.00176 | | #N/A | #N/A | 0.00051 | #N/A | W/W# | | 0.01119 | | 0.03205 | 0.01753 | | 0.00003 | |
| III IIIcras | #N/A | 7.08 | #N/A | 10.42 | | 6.55 | | #N/A | Δ/N# | 4 18 | W/V# | W/N# | | 3.63 | | 7.79 | 22.89 | | 4.14 | |
| KIAA0175 gene | product | integral type I protein | EST | calponin 2 | trinucleotide repeat containing 11 (THR-associated protein, | 230 kDa subunit) | minichromosome maintenance deficient | (mis5, S. pombe) 6 | zinc finger protein 142 | villin-like | EST | KIAA0440 protein | replication factor C | (activator 1) 4 (37kD) | Neil blood group precursor (McLeod | phenotype) | KIAA0715 protein procollagen-proline, 2- | oxoglutarate 4- dioxygenase (proline 4- | hydroxylase), alpha polypeptide II | - |
| n bac | | | 1738 | 1747 | | 1748 | • | 1749 | | | | 1784 | | 1786 | | 1791 | 1803 | | 1808 | 1 |
| Сепрапк | D79997 | D80710 | D80948 | D83735 | | D83783 | | D84557 | 570780 | D88154 | E02330 | F03811 | | F04320 | | F04531 | F09394 | | F09788 | |
| Arry ID | D79997 at | rc D80710 f at | rc D80948 f at | D83735 at | I | D83783_at | | D84557_at | . \$0.020 | D88154 at | 200134 at | rc F03811 f at | | rc_F04320_s_at | | rc F04531 s at | rc_F09394_s_at | | rc F09788 at | |
| | Arry ID Genbank Seq ID Known Gene Name III metas pivalue mean mean mean mean mean mean mean mea | Cenbank Seq ID Known Gene Name III litetas p value Integral Media Car Develor Integral Media Car Develor Integral Media Car Develor III litetas p value Integral Media Car Develor III litetas p value Integral Media Car Develor III litetas p value | Genbank Seq ID Known Gene Name III litetas p Value Inferial Media Car Development Car Developm | Genbank Seq ID Known Gene Name Interest Plant Model KIAA0175 gene KIAA0175 gene #N/A #N/A 55.25 42.49 53.87 8.57 6.15 at D80710 1734 integral type I protein 7.08 0.00213 253.6 303.7 149.08 22.95 28.29 at D80948 1738 EST #N/A #N/A #N/A 82.4 67.39 42.83 41.25 43.53 | ACCENDANCE Seq 1D NATIONAL Gene HIGH SECTION CONTROLL SECTION CONTROLL SECTION CONTROLL SECTION CONTROLL SECTION CONTROLL SECTION CONTROLL SECTION CONTROL CON | KIAA0175 gene | Companies Comp | KiAA0175 gene | KIAA0175 gene | Company Seq ID National Center Native III Interest places March III Interest Mar | Kindle K | Canbank Seq ID Nation Cerie Native Interest placed interests placed interests placed interests placed interests placed interests KIAA0175 gene HNIA HNIA 55.25 42.49 53.87 8.57 6.15 28.29 | KIAAO175 gene | Carindank Sag ID Anown Cente Marine In Instate prattice in Fortiary (Cente) Cente) Cente Cente) Cente Cente) Cente Ce | Contidency Con | Combanic Seq 10 National British Section 17:08 Control Ordinary Combanic Section 17:08 Control Ordinary Control Ordin | Drawing Seq ID National Carlos Native Integral (NAMO) To gene | DR37997 1725 product Milk Milk Milk 55.25 42.49 53.87 8.57 6.15 | Compared Seq Data National Control C | Contidant Seq 10 Contident S |

Table 9A. Genes and ESTs expressed only in metastatic liver turnor versus normal sample set 2.

| | | | | told | | | | | normal | normal | normal |
|----------------|---------|--------|---|----------|---------|-------------------------|---------|-------------|---------|---------|---------|
| | | | | change | - | metastatic: metastatic: | | metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| rc_H03629_s_at | H03629 | 1834 | desmin | W/A# | #N/A | 52.19 | 58.52 | 35.47 | -1.96 | -2.94 | 8.58 |
| rc_H09281_at | H09281 | 1863 | EST | 9 | 0.00966 | 147.11 | 187.25 | 71.67 | -2.97 | -0.54 | 11.7 |
| rc_H11760_f_at | H11760 | 1878 | EST | #N/A | #N/A | . 70.59 | 62.53 | 58.16 | 96.6 | 8.19 | 11.54 |
| | | | E74-like factor 4 (ets | | | | | | | | |
| | | | domain transcription | | | | • | | | | |
| rc_H24269_s_at | H24269 | 1902 | factor) | #N/A | #N/A | 51.79 | 47.58 | 24.25 | 15.19 | 16.8 | 12.36 |
| | | | gamma-aminobutyric acid (GABA) B | | | | | | | | |
| H43286_s_at | H43286 | 1929 | receptor, 1 | 5.05 | 0.01972 | 377.04 | 222.91 | 315.96 | 57.02 | 63.58 | 24.25 |
| | | | H2A histone family, | | | | | | | | |
| rc_H43646_at | H43646 | 1930 | member Y | 4.6 | 0.00147 | 413.02 | 424.34 | 219.27 | 92.12 | 91.3 | 52.41 |
| H53657_s_at | H53657 | 1945 | adenylate cyclase 3 | 3.98 | 0.0045 | 140.07 | 111.99 | 74.77 | 31.33 | 31.74 | 15.29 |
| rc_H57709_s_at | H57709 | 1956 | ribosomal protein L31 | #N/A | #N/A | 47.56 | 34.17 | 56.68 | 4.55 | 2.44 | 52.46 |
| | | | solute carrier family 2 | | | | | | | | |
| | | | (racilitated glucose transporter), member | | | | | | | | |
| rc_H58873_s_at | H58873 | 1961 | | 57.98 | 0.00063 | 4996.66 | 4603.55 | 4107.24 | 69.89 | 70.74 | 58.94 |
| rc_H59617_at | H59617 | 1964 | EST | 3.3 | 0.04588 | 116.07 | 71.39 | 101 | 19.24 | 21.65 | 16.88 |
| rc_H78211_at | H78211 | 2001 | EST | 6.73 | 0.02488 | 211.51 | 183.81 | 208.45 | -115.95 | -129.18 | 70.22 |
| rc_H98657_at | H98657 | 2068 | EST | #N/A | #N/A | 179.15 | 192.17 | 32.25 | 111.2 | 84.74 | 128.92 |
| | | • | | | | | | | | | |
| | | | creatine kinase, | | , | | | | | | • |
| | 00110 | | mitochondrial 1 | 1 | . (| 0 | | 0,000 | į | | 0 |
| J04469_at | J04469 | 2111 | (ubiquitous) | Ð./ | 0.00705 | 212.04 | 143.89 | 162.12 | -17.21 | -18.81 | 12.68 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| 1 |
|--|
| Genbank Seg ID Known Gene Name |
| 1 |
| 2119 dehydrogenase 1 mucin 1, |
| 2121 transmembrane ribosomal protein S6 kinase, 90kD, |
| 2146 polypeptide 1 trefoil factor 3 |
| 2149 (intestinal) tetracycline |
| 2157 transporter-like protein matrix metalloproteinase 12 (macrophage |
| 2179 elastase) ribose 5-phosphate isomerase A (ribose 5- |
| 2201 phosphate epimerase) endothelial cell protein C/activated protein C |
| |
| 2208 (autoantigenic) protease, serin |
| 2214 (prostasin) |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | plol | | | | | normal | normal | normal |
|-------------|---------|--------|--|----------|---------|-------------|-------------------------|-------------|--------|---------|---------|
| | | | • | change | | metastatic: | metastatic: metastatic: | metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Seq ID Known Gene Name | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| L42583_f_at | L42583 | 2215 | keratin 6A | #N/A | #N/A | 116.33 | 111.15 | 91.3 | 60.92 | 51.12 | 22.52 |
| M12125_at | M12125 | 2241 | tropomyosin 2 (beta) | 10.83 | 0.00191 | 291.63 | 158.55 | 229.08 | 13.01 | 16.33 | 11.64 |
| | | | thymidine kinase 1, | , | ٠. | | | | | | |
| M15205_at | M15205 | 2265 | soluble | 3.75 | 0.00159 | 153.15 | 150.31 | 66.27 | 41.98 | 35.02 | 29.02 |
| M16364_s_at | M16364 | 2269 | creatine kinase, brain | 12.69 | 0.03633 | 683.38 | 491.19 | 929.94 | -72.18 | -70.9 | 40.37 |
| l t | | | carcinoembryonic | | | | | | | | |
| · | | | antigen-related cell | | | | , | | | | |
| | | | adhesion molecule 6 | | | | | | - | | |
| | | | (non-specific cross | | | | | | | | |
| M18728_at | M18728 | 2285 | reacting antigen) | 44.82 | 0.00291 | 1390.62 | 1362.73 | 1031.01 | -9.17 | -13.17 | 11.83 |
| | | | renal/pancreas/salivar | | | | | | | | |
| M25629_at | M25629 | 2307 | · | #N/A | #N/A | 47.43 | 40.22 | 29.15 | -2.42 | 2.12 | 15.6 |
| | • | | endogenous retroviral | | | | | | | | |
| M27826_at | M27826 | 2313 | protease | 26.36 | 0.00342 | 993.89 | 833.56 | 816.33 | 5.88 | 7.04 | 32.2 |
| | | | melanoma adhesion | | | | | | | | |
| M29277_at | M29277 | 2316 | molecule | 3.91 | 0.00112 | 269.54 | 313.13 | 82.76 | 80.09 | 81.21 | 48.32 |
| | | | carcinoembryonic | | | | | | | | |
| | | | antigen-related cell | | | | | | | | |
| M29540_at | M29540 | 2317 | adhesion molecule 5 | 36.57 | 0.0116 | 1516.55 | 1083.69 | 1372.55 | -1.09 | -3.15 | 11.75 |
| | | | transcription factor 3 (E2A immunoglobulin | | | | | | | | |
| | | | enhancer binding | | | | | | | | |
| M31523_at | M31523 | 2329 | factors E12/E47) | #N/A | #N/A | 59.78 | 62.53 | 10.63 | 18.65 | 21.63 | 8.94 |
| M87230 of | 1497330 | 2445 | replication factor C | 707 | 0.00348 | 02 72 | 70.4 | 92 | 96 7 | 0 | 16 70 |
| מוס נסמי | 200 | 217 | (activated 1) + (5) hz/) midkine (neurite | ř | 2000 | 37.76 | - 6 | 90.00 | 4.20 | 76.6 | 0.0 |
| | | · | growth-promoting | | • | | | | | | |
| M94250_at | M94250 | 2426 | factor 2) | 10.39 | 0.01818 | 442.2 | 188.32 | 624.9 | -155.4 | -175.43 | 101.89 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | fold | | | | | normal | normal | normal |
|----------------|---------|--------|---|----------|---------|-------------|-------------|-------------|--------|--------|---------|
| | | | | change | - | metastatic: | metastatic: | metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| M04345 of | M94345 | 2427 | capping protein (actin | 22.38 | 0.00003 | 503 47 | 539 15 | 179 05 | -7.51 | -10.35 | 27 75 |
| M94362_at | M94362 | 2428 | lamin B2 | #N/A | #N/A | 175.16 | 154.32 | 75.52 | 79.2 | 81.73 | 34.64 |
| I | | | pregnancy specific beta-1-glycoprotein 4 pregnancy specific | | | | | | | | |
| M94891_s_at | M94891 | 2429 | beta-1-glycoprotein 7 hydroxymethylbilane | #N/A | #N/A | 221.09 | 197.74 | 80.32 | 178.23 | 166.98 | 61.33 |
| M95623 cds1 at | M95623 | 2431 | synthase | #N/A | #N/A | 67.65 | 72.44 | 22.29 | 6.67 | 0.88 | 24.92 |
| rc_N22015_at | N22015 | 2448 | EST | 46.61 | 0.00025 | 1225.51 | 887.65 | 1106.3 | -5.3 | -6.84 | 18.82 |
| rc_N35376_at | N35376 | 2501 | EST | #N/A | #N/A | 39.01 | 35.99 | 19.81 | 28.53 | 29.44 | 8.2 |
| | | | v-myb avian | | | | | | | | |
| rc N49284 s af | N49284 | 2537 | iniyeloblastosis viral oncodene homolog | 11.82 | 0.01981 | 510.82 | 523.45 | 423.41 | -36.17 | -50.08 | 50.71 |
| rc N54265 s at | N54265 | 2563 | EST | #N/A | W/A | 45.56 | 44.98 | 19.91 | 17.71 | 26.43 | 25.36 |
| rc N54395 at | N54395 | 2565 | EST | #N/A | #N/A | 49.95 | 46.84 | 48.26 | -16.14 | -17.71 | 6.65 |
| I | | | DEAD/H (Asp-Glu-Ala- Asp/His) box | | | | | | | | |
| rc_N62675_s_at | N62675 | 2594 | polypeptide 16 | 3.61 | 0.04034 | 109.44 | 104.65 | 108.08 | 6.49 | 15.91 | 35.18 |
| rc N63165 at | N63165 | | EST | #N/A | #N/A | 68.04 | 62.1 | 44.7 | 30.07 | 26.23 | 13.84 |
| rc_N64616_at | N64616 | | EST | 3.11 | 0.0074 | 68.89 | 75.8 | 32.12 | -0.83 | 9.77 | 28.69 |
| rc_N66951_at | N66951 | 2621 | EST | 5.54 | 0.02442 | 451.16 | 432.45 | 381.81 | 59.83 | 75.27 | 32.75 |
| | | | solute carrier family 11 (proton-coupled divalent metal ion | | | | | | | | |
| rc N72116 s at | N72116 | 2668 | transporters), member 2 | 9 04 | 0 00051 | 338 43 | 314.06 | 196.66 | 32.57 | 29.68 | 20.1 |
| rc_N73846_at | N73846 | 2680 | EST | 3.27 | 0.00012 | 78.37 | 79.7 | 20.06 | 14.43 | 14.3 | 17.21 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| • | | | | | fold | | | | | normal | normal | normal |
|---|----------------|---------|--------|---|----------|---------|-------------|-------------------------------------|-------------|--------|--------|---------|
| | | | | | change | _ | netastatic: | metastatic: metastatic: metastatic: | metastatic: | set 2: | set 2: | set 2: |
| | Affy ID | Genbank | Sed ID | Known Gene Name | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| • | rc N89670 at | 02968N | 2709 | | 4.26 | 0.00002 | 115.98 | 108.4 | 29.79 | 3.16 | -1.81 | 43.32 |
| | rc_N92659_at | N92659 | 2721 | EST | #N/A | #N/A | 31.19 | 37.23 | 14.14 | 5.91 | 4.05 | 5.63 |
| , | rc_N92734_at | N92734 | 2722 | EST | #N/A | #N/A | 45.63 | 39.49 | 14.19 | 27.31 | 25.45 | 10.41 |
| | i i | | | protein tyrosine | | | | | | | | |
| | | | | phosphatase type IVA, | | | | | | | | |
| | rc_N93798_at | N93798 | 2738 | member 3 | 4.65 | 0.00118 | 557.51 | 523.56 | 229.03 | 139.69 | 122.61 | 84.93 |
| | i I | | | tumor protein D52-like | | | | | | | | |
| | rc R06251 f at | R06251 | 2764 | 2 | 5.57 | 0.00037 | 343.86 | 381.79 | 95.11 | 74.25 | 66.72 | 64.11 |
| | rc R06866 s at | R06866 | 2774 | EST | 5.18 | 0.00187 | 148.48 | 114.86 | 102.01 | 18.26 | 15.16 | 17.11 |
| | i i | | | midline 1 (Opitz/BBB | | | | | | | | |
| | rc_R26744_at | R26744 | 2804 | syndrome) | 4.32 | 0.00532 | 112.54 | 06 | 76.53 | -0.2 | -3.06 | 24.42 |
| | rc_R27296 f at | R27296 | 2807 | EST | #N/A | #N/A | 41.82 | 39.83 | 32.72 | 8.27 | 11.07 | 7.26 |
| | rc_R33498_s_at | R33498 | 2820 | EST | 41.34 | 0.00001 | 1839.74 | 1920.41 | 1082.84 | 46.45 | 33.01 | 43.64 |
| | rc_R36109_at | R36109 | 2823 | EST | #N/A | #N/A | 47.54 | 48.43 | 19.9 | 25.67 | 20.25 | 34.73 |
| | į | | | calcium channel, | | | | | | | | |
| | | ! | 1 | voltage-dependent, | • | | | | - | | | 1 |
| | rc_R36947_s_at | R36947 | 2825 | beta 3 subunit protein similar to E.coli | 4.11 | 0.00006 | 101.82 | 109.62 | 78.57 | -0.61 | 0.94 | 37.6 |
| | | | | yhdg and R. | | | | | | | | |
| | rc_R38511_s_at | R38511 | 2832 | capsulatus nifR3 | 5.19 | 0.00015 | 131.5 | 147.54 | 44.02 | 21.3 | 23.75 | 9.22 |
| | rc_R39191_s_at | R39191 | 2834 | KIAA1020 protein | 4.69 | 0.00456 | 130.93 | 145.31 | 62.13 | 17.46 | 18.33 | 16.04 |
| | | | | prominin (mouse)-like | | | | | | | | |
| | rc_R40057_at | R40057 | 2839 | | #N/A | #N/A | 47.6 | 46.28 | 36.57 | 0.83 | -0.8 | 7.91 |
| | | | | KIAA0552 gene | | | | | | | | |
| | rc_R44479_at | R44479 | 2855 | product | 4.14 | 0.0181 | 97.01 | 105.95 | 60.51 | 7.08 | 5.99 | 7.62 |
| | rc_R44817_at | R44817 | 2860 | EST | #N/A | #N/A | 600.1 | 550.42 | 309.14 | 255.91 | 249.29 | 60.69 |
| - | | | | Fc fragment of IgG, low affinity IIIa. | | | | | | | | , |
| | rc_R49047_at | R49047 | 2878 | receptor for (CD16) | #N/A | #N/A | 44.49 | 36.71 | 14.37 | 15.43 | 14.16 | 14.13 |
| | i | | | | | | | | | | | |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | FOIN | | | | | normal | normal | norma |
|----------------|---------|--------|---|----------|---------|-------------|-------------------------|-------------|---------|---------|---------|
| | | | | 2000 | | motoctatic. | motactatic: motactatic: | metactatic. | sof 2. | set 2. | set 2. |
| Affv ID | Genbank | Sea 1D | Genbank Sed ID Known Gene Name | in metas | p value | Mean | Median | | Mean | Median | Std Dev |
| | | | ubiquitin specific | | | | | | | | |
| | | | protease 7 (herpes | | | | | | | | |
| rc R54935 i at | R54935 | 2903 | virus-associated) | #N/A | #N/A | 75.76 | 81.15 | 31.57 | 17.77 | 14 | 18.86 |
| rc R55470 at | R55470 | 2904 | EST | 3.59 | 0.00515 | 256.38 | 225.69 | 138.22 | 75.29 | 73.42 | 53.04 |
| R56678 at | R56678 | 2908 | EST | 3.81 | 0.02242 | 98.37 | 78.38 | 75.74 | 2.67 | ß | 5.82 |
| rc R59093 at | R59093 | 2911 | EST | #N/A | #N/A | 57.79 | 33.48 | 73.58 | 12.26 | 10.21 | 7.73 |
| rc_R63925_at | R63925 | 2929 | EST | #N/A | #N/A | 70.52 | 60.93 | 15.36 | 49.82 | 52.73 | 13.77 |
| R69700 at | R69700 | 2943 | EST | 6.71 | 0.0021 | 387.34 | 393.81 | 94.86 | 79.33 | 57.85 | 129.97 |
| rc R71395 at | R71395 | 2952 | EST | 10.42 | 0.00422 | 318.75 | 274.93 | 227.75 | 14.99 | 12.28 | 14.58 |
| R76363 at | R76363 | 2962 | EST | #N/A | #N/A | 47.81 | 38.04 | 24.76 | 12.67 | 16.94 | 13.17 |
| rc R85266 at | R85266 | 2977 | EST | #N/A | #N/A | 53.4 | 54.53 | 25.44 | 48.68 | 39.83 | 33.9 |
| rc_R91819_at | R91819 | 2984 | EST | 8.95 | 0.0000 | 263.33 | 219.91 | 135.67 | 11.34 | 9.35 | 36.49 |
| l I | | | matrix | | | | | | | | |
| | | | metalloproteinase 12 | | | | | | | | |
| | | | (macrophage | | | | | | | | |
| rc R92994 s at | R92994 | 2990 | elastase) | 11.05 | 0.00248 | 312.14 | 252.62 | 248.32 | 11.43 | 6.64 | 11.52 |
| rc R95966 i at | R95966 | 2997 | EST | 11.22 | 0.00682 | 482.68 | 436.3 | 446.39 | -106.64 | -160.75 | 127.91 |
| rc_R96924_s_at | R96924 | 3001 | EST | 6.18 | 0.03417 | 451.59 | 490.51 | 339.66 | 51.01 | 52.94 | 54.09 |
| S78187 at | S78187 | 3036 | cell division cycle 25B | 8.07 | 0.0000 | 198.51 | 218.04 | 74.52 | 3.36 | -1.2 | 26.9 |
| rc T03438 s at | T03438 | 3043 | EST | 8.18 | 0.00032 | 300.09 | 229.6 | 218.02 | 31.03 | 28.88 | 15.86 |
| | T03541 | 3045 | EST | #N/A | #N/A | 455.39 | 418.21 | 288.45 | 155.27 | 155.62 | 50.93 |
| rc_T15473_at | T15473 | 3058 | muscle specific gene | 5.81 | 0.02404 | 189.25 | 139.11 | 184.39 | -5.65 | -10.77 | 15.89 |
| | | | cleavage and | | | | | | | | |
| | | | polyadenylation specific factor 4, 30kD | | | | | | | | |
| rc T16983 s at | T16983 | 3074 | subunit | 5.23 | 0.00075 | 268.21 | 300.53 | 81.43 | 65.64 | 45.67 | 74.28 |
| rc_T25744_s_at | T25744 | 3092 | EST | #N/A | #N/A | 79.78 | 68.72 | 52.01 | 15.56 | 12.65 | 13.18 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | - | | | fold | | | | | normal | normal | normal |
|----------------|---------|------|------------------------------|----------|---------|-------------|-------------|-------------------------------------|--------|--------|---------|
| | | | | change | - | metastatic: | metastatic: | metastatic: metastatic: metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | | Seq ID Known Gene Name | In metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | protease, serine, 8 | | | | - | | | | |
| rc_T30193_s_at | T30193 | 3098 | (prostasin) | 8.39 | 0.00043 | 1912.22 | 1228.46 | 1739.5 | 214.88 | 175.57 | 127.28 |
| rc T30222 at | T30222 | 3100 | EST | W/A# | #N/A | 35.41 | 34.75 | 27.14 | 8.47 | 9.56 | 17.71 |
| rc_T32108_at | T32108 | 3103 | EST | 96.9 | 0.00723 | 1095.16 | 593.52 | 923.47 | 148.09 | 152.11 | 92.06 |
| rc_T47601_at | T47601 | 3126 | EST | 4.05 | 0.00878 | 199.79 | 219.84 | 62.16 | 84.5 | 28.33 | 122.16 |
| rc T53404 at | T53404 | 3143 | EST | 10.68 | 0.00582 | 654.13 | 475.25 | 687.49 | 48.3 | 9.0 | 93.06 |
| rc_T66935_at | T66935 | 3179 | EST | 3.97 | 0.00188 | 253.93 | 230.32 | 129.66 | 8.99 | 52.94 | 42.32 |
| rc_T89601_r_at | T89601 | 3243 | EST | #N/A | #N/A | 839.07 | 747.51 | 231.92 | 410.07 | 342.22 | 200.88 |
| rc T91116 at | T91116 | 3252 | EST | 4.01 | 0.02721 | 133.54 | 61.72 | 126.12 | 16.62 | 11.42 | 13.64 |
| rc_T96060_at | T96060 | 3263 | EST | #N/A | #N/A | 910.76 | 225.75 | 1282.67 | 67.92 | 70.44 | 91.72 |
| l I | | | inositol 1,4,5- | | | • | | | | | |
| ٠ | | | triphosphate receptor, | • | | | | | | | |
| U01062_at | U01062 | 3273 | type 3 | 7.41 | 0 | 160.46 | 164.36 | 23.91 | -16.8 | -26.61 | 26.44 |
| , | | | active BCR-related | | | | | | | | |
| U01147_at | U01147 | 3275 | gene | 3.22 | 0.00103 | 97.43 | 84.12 | 41.43 | 27.15 | 22.94 | 17 |
| | | | tumor necrosis factor | | | | | | | | |
| | | | (ligand) superfamily, | | | | | | | | |
| U03398_at | U03398 | 3282 | member 9 | #N/A | #N/A | 116.17 | 88.07 | 59.5 | 115.08 | 92.7 | 93.14 |
| | | | protease inhibitor 5 | | | | | | | | |
| U04313_at | U04313 | 3284 | (maspin) | 4.54 | 0.02986 | 132.77 | 69.67 | 115.99 | -0.34 | -2.59 | 9.7 |
| | | | cadherin 17, LI | | | | | 4 | | | |
| | | | cadherin (liver- | | | | | | | • | |
| U07969_s_at | O07969 | 3289 | intestine) | 10.78 | 0.02002 | 428.65 | 383.68 | 390.23 | 15.78 | 12.82 | 12.06 |
| | | | chondroitin sulfate | | | | | | | | |
| U16306 at | U16306 | 3312 | proteogrycan z (versican) | W/N# | #N/A | . 48.35 | 56.91 | 35.99 | 8.07 | 4.58 | 16.58 |
| 5 | | | | , | | | | | | | |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | fold | | | | | normal | normal | normal |
|------------------------|------------------|--------------|--|--------------|-----------------|-------------|-------------------------|----------------|----------------|----------------|---------------|
| | | | | change | _ | metastatic: | metastatic: metastatic: | metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | Mean | | | Mean | Median | Std Dev |
| | | | laminin, beta 3 (nicein (125kD), kalinin (140kD), BM600 | | | | | | | | |
| U17760_rna1_at | U17760 | 3315 | (125kD)) sulfotransferase family | 3.54 | 0.01853 | 103.13 | 111.96 | 72.83 | 6.99 | -3.15 | 21.08 |
| U20499_at | U20499 | 3321 | 1A, phenol-preferring, member 3 | 5.5 | 0.00299 | 316.7 | 231.67 | 222.02 | 48.34 | 54.37 | 24.69 |
| | | | epithelial protein up- regulated in carcinoma, membrane | | | | | | | | |
| U21049_at | U21049 | 3325 | associated protein 17 TAR (HIV) RNA- | 7.53 | 0.01667 | 202.38 | 248.15 | 119.31 | -14.32 | -12.21 | 19.43 |
| U38847_at | U38847 | 3357 | binding protein 1 potassium voltage- gated channel, KQT- | #N/A | #N/# | 72.74 | 66.76 | 36.14 | 15.7 | 17.43 | 10.2 |
| U40990_at | U40990 | 3359 | like subfamily, member 1 discoidin domain | 3.18 | 0.00093 | 128.02 | 142.31 | 43.16 | 40.88 | 42.64 | 23.09 |
| U48705_rna1_s_at | U48705 | 3370 | receptor family, member 1 caudal type homeo | 5.94 | 0.01323 | 178.04 | 200.63 | 102.82 | -1.87 | -6.25 | 25.81 |
| U51095_at U53786_at | U51095 U53786 | 3382 3390 | factor 1 envoplakin gamma-aminobutyric | 4.76 #N/A | 0.02664 #N/A | 130.81 | 143.82 73.95 | 93.4 249.37 | 5.83 -20.54 | 5.55 -20.08 | 8.97 16.22 |
| U66661_at | U66661 | . 3406 | acid (GABA) A receptor, epsilon | #N/A | #N/A | 55.33 | 44.93 | 29.26 | 16.58 | 14.81 | 11.96 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | fold | | | | | norma | normal | norma |
|----------------|------------------|--------|---|--------------|---------|-------------|-------------------------------------|-----------------|--------|--------|---------------|
| | | | | | | | | | 3 | | |
| | | | | cnange | | metastatic: | metastatic: metastatic: metastatic: | metastatic: | set z: | set z: | :7 18S |
| Affy ID | Genbank | Sed ID | | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | DEAD/H (Asp-Glu-Ala- Asp/His) box polypeptide 11 (S.cerevisiae CHL1- | | | | | | | | |
| U75968_s_at | U75968 | 3423 | like helicase) O-linked N- | #N/A | #N/A | 195.48 | 155.37 | 89.8 | 115.5 | 102.24 | 77.72 |
| | | | acetylglucosamine (GlcNAc) transferase | | | | | | | | |
| | | | (UDP-N- | | | | | | | | |
| ٠ | | | ypeptide-N- | | | | | | | | |
| U77413_at | U77413 | 3427 | acety/glucosaminyl ransferase) | #N/A | #N/A | 59.38 | 59.78 | 48.05 | 1.2 | 5.23 | 13.8 |
| ** 000001 | 00001 | 0.450 | pyridoxal (pyridoxine, | . 64 | 66600 | | 0.4 | 1000 | 7 7 | 0,00 | , 1 |
| U69606_at | 009000 W02605 | 3452 | VICALIIIII DO) NIIIASE | 5.30 #N/₩ | 4N/A | 120 58 | 97.30 | 36.63 100 52 | 30.50 | 33.06 | 74.3 16.14 |
| 18_C2030_ | 1102030 | 5 | | | | 20.00 | 30.06 | 100.02 | 9 | 9 | 5 |
| | | | leukemia innibitory factor (cholinergic | ٠ | | | | | | | |
| rc_W46451_s_at | W46451 | 3529 | differentiation factor) | #N/A | #N/A | 120.95 | 76.95 | 88.42 | 36.63 | 40.09 | 21.06 |
| rc_W60968_at | W60968 | 3559 | EST | #N/A | #N/A | 125.7 | 144.72 | 48.22 | 51.99 | 55.29 | 21.11 |
| rc_W67251_s_at | W67251 | 3570 | EST | 6.13 | 0.01463 | 204.71 | 182.17 | 127.82 | 21.77 | 23.88 | 12.17 |
| rc_W73189_at | W73189 | 3589 | EphB2 | 3.69 | 0.02909 | 113.63 | 144.75 | 67.73 | 20.7 | 23.2 | 15.44 |
| rc_W78057_at | W78057 | 3600 | EST | 90.6 | 0.0034 | 397.29 | 374.78 | 305.93 | 29.21 | 29.9 | 34.33 |
| rc_W90146_f_at | W90146 | 3644 | EST | 6.23 | 0.01558 | 170.66 | 147.78 | 126.32 | 9.93 | 8.63 | 6.49 |
| rc_W92449_at | W92449 | 3652 | EST | 31.67 | 0.00011 | 715.17 | 491.5 | 459.71 | -40.13 | -40.74 | 17.76 |
| | | | protease inhibitor 5 | | | | | | | | |
| rc_W93726_s_at | W93726 | 3656 | (maspin) | 16.48 | 0.00014 | 355.41 | 304.26 | 149.69 | -14.2 | -14.8 | 10.59 |
| W95348_at | W95348 | 3663 | HSPC113 protein | 10.89 | 0.01065 | 555.52 | 492.63 | 563.86 | 26.59 | 29.36 | 21.03 |
| rc_W95477_at | W95477 | 3664 | EST | 26.51 | 0.00161 | 941.08 | 9.999 | 1130.33 | 17.15 | 18.75 | 12.83 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| | | | | fold | | | | | normal | normal | normal |
|-------------|---------|--------|-------------------------|----------|---------|-------------|-------------------------|-------------|--------|--------|---------|
| | | | | change | | metastatic: | metastatic: metastatic: | metastatic: | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| | | | small nuclear | | | | | | | | |
| | | | ribonucleoprotein | | | | | | | | |
| | | | 70kD polypeptide | | | | | | | | ! |
| X04654_s_at | X04654 | 3681 | (RNP antigen) | #N/A | #N/A | 98.11 | 89.35 | 38.15 | 42.22 | 41.24 | 17.18 |
| X13956 at | X13956 | 3701 | EST | 3.2 | 0.00321 | 79.19 | 75.13 | 39.53 | 15.11 | 18.02 | 14.55 |
| l | | | H2A histone family, | | | | | | | | |
| X14850_at | X14850 | 3706 | member X | 4.11 | 0.0001 | 118.41 | 97.53 | 52.6 | 11.6 | 13.26 | 30.46 |
| | | | v-abl Abelson murine | | | | | | | | |
| | | | leukemia viral | | | | | | | | |
| X16416_at | X16416 | 3713 | oncogene homolog 1 | #N/A | #N/A | 82.49 | 90.74 | 16.7 | 36.59 | 38 | 14.59 |
| | i i | i | | 0 | 0 | 0 | 7000 | 177 | 0 | U 7 | 74 00 |
| X54667_s_at | X5466/ | 3/31 | cystatin S, cystatin SN | 8.53 | 0.00038 | 2/3.90 | 109.84 | 11./12 | -10.09 | -13.00 | 4.03 |
| X57348 s at | X57348 | 3744 | stratifin | 12.53 | 0.0013 | 308.28 | 241.69 | 194.79 | -63.66 | -76.43 | 44.95 |
| l İ | | | matrix | | | | | | | | |
| | | | metalloproteinase 11 | | | | | | | | |
| X57766_at | X57766 | 3745 | (stromelysin 3) | #N/A | #N/A | 166.25 | 142.96 | 124.34 | 63.38 | 62.13 | 25.48 |
| 1 | | | cadherin 3, P-cadherin | | | | | | | | |
| X63629_at | X63629 | 3762 | (placental) | 3.02 | 0.01654 | 67.22 | 76.67 | 29.17 | -4.24 | -6.82 | 16.9 |
| | | | interferon, alpha- | | | | | | | | |
| X67325_at | X67325 | 3775 | inducible protein 27 | 9.67 | 0.03245 | 962.87 | 412.22 | 1361.55 | 26.81 | 48.73 | 69.77 |
| | | | sodium channel, | | | | | | | | |
| | | | nonvoltage-gated 1 | | | | | | | • | |
| X76180 at | X76180 | 3795 | alpha | 11.68 | 0 | 320.05 | 268.84 | 127.86 | 22.38 | 23.9 | 15.02 |
| 1 | | | cadherin 17, LI | | | | | | | | |
| | | | cadherin (liver- | | | | | | | | |
| X83228 at | X83228 | 3810 | intestine) | 10.58 | 0.02147 | 342.12 | 423.87 | 282.49 | -8.87 | 6.9- | 8.55 |
| I | | | FXYD domain- | | | | | | | | |
| | | , | containing ion | | 1 | | | 9 | 6 | L | 9 |
| X93036_at | X93036 | 3830 | transport regulator 3 | 42.36 | 0.00167 | 1322.91 | 783.61 | 1219 | -83.87 | -85.84 | 40.53 |

Table 9A. Genes and ESTs expressed only in metastatic liver tumor versus normal sample set 2.

| Genbank Seq ID Known Gene Name in metas p value Mean Median S X99133 3842 24p3) 6.27 0.0453 284,56 129,1 4 Y00503 3849 keratin 19 14.19 0.00217 362.5 427.85 2 HMT1 (hnRNP methyltransferase, S. 4.28 0.00124 392.5 427.85 2 it Z39191 3901 EST #N/A #N/A 166.55 125.78 2 it Z39569 3909 EST #N/A #N/A 166.55 125.78 2 SRY (sex-determining region Y)-box 9 (campomelic dysplasia, autosomal (campomelic dysplasia, autosomal #N/A #N/A 39.69 40.61 | | | | | fold | | | | | norma | normal | normal |
|--|--------------|---------|--------|--------------------------------------|----------|---------|-------------|-------------|---------|--------|--------|---------|
| Genbank Seq ID Known Gene Name in metas p value Mean Median X99133 3842 24p3) 6.27 0.0453 284.56 129.1 Y00503 3849 keratin 19 14.19 0.00217 362.5 427.85 HMT1 (hnRNP methyltransferase, S. 4.28 0.00124 393.27 449.97 Z39191 3901 EST #N/A #N/A #N/A 166.55 125.78 Z39569 3909 EST #N/A #N/A 199.73 88.96 Z41415 3933 EST #N/A #N/A 199.73 88.96 Campomelic dysplasia, autosomal #N/A #N/A #N/A 39.69 40.61 | | | | • | change | | metastatic: | metastatic: | | set 2: | set 2: | set 2: |
| Secondary Seco | Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | Mean | Median | Std Dev | Mean | Median | Std Dev |
| X99133 3842 24p3) 6.27 0.0453 284.56 129.1 Y00503 3849 keratin 19 14.19 0.00217 362.5 427.85 HMT1 (hnRNP methyltransferase, S. methyltransferase, S. 4.28 0.00124 393.27 449.97 Z39191 3901 EST #N/A #N/A #N/A 166.55 125.78 Z41415 3933 EST #N/A #N/A 199.73 88.96 Campomelic campomelic dysplasia, autosomal #N/A #N/A 39.69 40.61 | | | | lipocalin 2 (oncogene | | | | | | | | |
| Y00503 3849 keratin 19 14.19 0.00217 362.5 427.85 HMT1 (hnRNP methyltransferase, S. methyltransferase, S. 4.28 0.00124 393.27 449.97 Z39191 3860 cerevisiae)-like 2 4.28 0.00011 442.36 371.88 Z39569 3909 EST #N/A #N/A #N/A 166.55 125.78 Z41415 3933 EST #N/A #N/A 199.73 88.96 Campomelic campomelic campomelic dysplasia, autosomal 40.61 Z46629 3938 sex-reversal) #N/A #N/A 40.61 | X99133_at | X99133 | 3842 | 24p3) | | 0.0453 | 284.56 | 129.1 | 434.01 | -28.39 | -19.41 | 26.24 |
| HMT1 (hnRNP methyltransferase, S. Y10807 3860 cerevisiae)-like 2 4.28 0.00124 393.27 449.97 239191 3901 EST 8.84 0.00011 442.36 371.88 239569 3909 EST #N/A #N/A 199.73 88.96 SRY (sex-determining region Y)-box 9 (campomelic dysplasia, autosomal 246629 3938 sex-reversal) #N/A #N/A 39.69 40.61 | Y00503_at | Y00503 | 3849 | keratin 19 | | 0.00217 | 362.5 | 427.85 | 240.73 | 7.02 | 10.74 | 10.98 |
| Methyltransferase, S. Y10807 3860 cerevisiae)-like 2 4.28 0.00124 393.27 449.97 Z39191 3901 EST 8.84 0.00011 442.36 371.88 Z39569 3909 EST #N/A #N/A 166.55 125.78 Z41415 3933 EST #N/A #N/A 199.73 88.96 SRY (sex-determining region Y)-box 9 (campomelic dysplasia, autosomal dysplasia, autosomal #N/A #N/A 39.69 40.61 | | | | HMT1 (hnRNP | | - | | | | | | • |
| Y10807 3860 cerevisiae)-like 2 4.28 0.00124 393.27 449.97 Z39191 3901 EST 8.84 0.00011 442.36 371.88 Z39569 3909 EST #N/A #N/A 166.55 125.78 Z41415 3933 EST #N/A #N/A 199.73 88.96 SRY (sex-determining region Y)-box 9 (campomelic dysplasia, autosomal dysplasia, autosomal #N/A #N/A 39.69 40.61 | | | | methyltransferase, S. | | | | | | | | |
| Z39191 3901 EST 8.84 0.00011 442.36 371.88 Z39569 3909 EST #N/A #N/A 166.55 125.78 Z41415 3933 EST #N/A #N/A 199.73 88.96 SRY (sex-determining region Y)-box 9 (campomelic dysplasia, autosomal 246629 3938 sex-reversal) #N/A #N/A 39.69 40.61 | Y10807_s_at | Y10807 | 3860 | cerevisiae)-like 2 | | 0.00124 | 393.27 | 449.97 | 142.94 | 96 | 90.42 | 44.52 |
| Z39569 3909 EST #N/A #N/A 166.55 125.78 Z41415 3933 EST #N/A #N/A 199.73 88.96 SRY (sex-determining region Y)-box 9 (campomelic dysplasia, autosomal 246629 3938 sex-reversal) #N/A #N/A 39.69 40.61 | rc_Z39191_at | Z39191 | 3901 | EST | 8.84 | 0.00011 | 442.36 | 371.88 | 228.18 | 46.98 | 49.3 | 33.31 |
| Z41415 3933 EST #N/A #N/A 199.73 88.96 SRY (sex-determining region Y)-box 9 (campomelic dysplasia, autosomal #N/A #N/A 39.69 40.61 | rc_Z39569_at | Z39569 | 3909 | EST | #N/A | #N/A | 166.55 | 125.78 | 217.77 | -9.57 | -12.15 | 17.32 |
| SRY (sex-determining region Y)-box 9 (campomelic dysplasia, autosomal #N/A #N/A 39.69 40.61 | rc Z41415 at | Z41415 | 3933 | EST | #N/A | #N/A | 199.73 | 88.96 | 271.2 | 10.38 | 6.64 | 11.47 |
| (campomelic dysplasia, autosomal Z46629 3938 sex-reversal) #N/A #N/A 39.69 40.61 | l I | | | SRY (sex-determining region Y)-box 9 | | | | | | | | |
| dysplasia, autosomal Z46629 3938 sex-reversal) #N/A #N/A 39.69 40.61 | | | | (campomelic | | | | | | | | |
| Z46629 3938 sex-reversal) #N/A #N/A 39.69 40.61 | | | | dysplasia, autosomal | | | | | | • | | |
| | Z46629_at | Z46629 | 3938 | sex-reversal) | #N/A | #N/A | 39.69 | 40.61 | 13.99 | 6.21 | 8.06 | 9.16 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affy ID | Genbank | Seq ID | Known Gene Name | fold change in metas | p value | metastati c: Mean | metastati metastati c: c: Std Median Dev | metastati c: Std Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|--|--|----------------------------|--|---|---|--|---|---|---|--|--|
| rc_AA001902_at rc_AA001903_i_at rc_AA004669_at | AA001902 AA001903 AA004669 | 9 0 0 | KIAA0305 gene product EST EST | #N/A #N/A #N/A | #N/A #N/A #N/A | 14.88 6.64 8.66 | 17.06 0.97 10.77 | 8.8 14.07 8.65 | 29.11 26.52 33.32 | 28.6 28.7 25.36 | 23.36 13.21 31.12 |
| rc_AA004707_at rc_AA005202_at | AA004707 AA005202 | 10 | copper chaperone for superoxide dismutase retinol-binding protein 4, interstitial | #N/A 3.18 | #N/A 0.00106 | 217.76 | 217.72 | 144.41 | 502.14 | 479.38 | 161.01 |
| rc_AA009719_at rc_AA010205_at rc_AA010360_at rc_AA010619_at | AA009719 AA010205 AA010360 AA010619 | 20 23 24 | peroxisomal membrane protein 2 (22kD) EST EST EST | 47.12 7.41 6.55 8.55 | 0.00008 0 0.00027 0.00057 | -50.14 14.43 12.5 21.58 | -51.69 17.64 14.77 5.47 | 47.17 14.87 8.62 42.99 | 1370.32 187.55 169.99 279.66 | 1503.99 154.99 135.98 268.6 | 715.62 92.13 129.2 154.98 |
| rc_AA013095_s_at rc_AA015768_at rc_AA016021_at rc_AA017146_at rc_AA018867_at rc_AA019715_at | AA013095 AA015768 AA016021 AA017146 AA018867 | 33 35 36 44 39 | potassium voltage-gated channel, shaker-related subfamily, beta member 1, EST ubiquitin-like 3 EST EST | #N/A 15.3 #N/A 10.1 42.87 #N/A | #N/A 0.00008 #N/A 0.00052 0.00002 #N/A | 7.81 12.22 13.88 30.35 45.29 8.03 | 3.06 14.2 16.21 22.85 29.28 8.03 | 19.71 11.71 13.21 50.05 52.68 6.49 | 18.56 417.95 65.65 414.24 1944.56 | 15.83 472.1 51.44 435.09 2160.33 | 8.24 248.15 46.44 193.39 1142.41 7.82 |
| rc_AA024511_at rc_AA024866_at | AA024511 AA024866 | 46 49 | suppressor of Ty (S.cerevisiae) 3 homolog EST | #N/A #N/A | #N/A #N/A | 7.82 | 3.08 | 12.06 8.56 | 62.54 31.16 | 58.34 30.56 | 27.73 14.26 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | norma | normal | normal |
|------------------|----------|--------|--|----------|----------|-----------|---------------------|---------------|--------|--------|---------|
| | • | | | opuedo | | motaetati | į | 740 | cot 2. | cat 2. | cot 2. |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median |) Oe Oe | Mean | Median | Std Dev |
| rc_AA025930_at | AA025930 | 52 | EST | 3.59 | 0.00372 | 33.24 | 25.62 | 26.11 | 115.2 | 110.13 | 45.17 |
| | | | | | | | | | | | |
| | | | microvascular endothelial | | | | | | | | |
| AA027766_at | AA027766 | 28 | differentiation gene 1 | #N/A | ¥N# | 16.71 | 16.37 | 7.36 | 25.05 | 26.19 | 13.26 |
| AA028976_at | AA028976 | 63 | EST | #N/A | #N/A | 9.46 | 10.28 | 18.91 | 72.06 | 47.28 | 55.89 |
| rc_AA031360_s_at | AA031360 | 29 | EST | #N/A | #N/A | 13.46 | 12.87 | 10.49 | 26.25 | 24.36 | 9.3 |
| rc_AA032250_at | AA032250 | 73 | EST | 3.56 | 0.000 | 10.56 | 13.86 | 9.55 | 84.44 | 83.69 | 50.06 |
| rc_AA034365_at | AA034365 | 9/ | EST | #N/A | #N/A | 119.56 | 39.46 | 171.27 | 216.28 | 159.86 | 152.39 |
| rc_AA039616_at | AA039616 | 06 | EST | 9.36 | 0.0000 | 2.48 | -4.83 | 14.2 | 238.71 | 224.42 | 125.28 |
| rc_AA040087_at | AA040087 | 92 | EST . | 4.13 | 0.00123 | 34.99 | 27.32 | 18.99 | 156.63 | 148.88 | 90.44 |
| rc_AA040291_at | AA040291 | . 46 | KIAA0669 gene product | 3.55 | 0.00308 | 12.98 | 12.72 | 16.36 | 101.69 | 98.94 | 80.43 |
| | | | | ` | | | | | | | |
| | * | | NADH dehydrogenase (ubiquinone) 1 beta | | | | | | | | |
| AA041208_at | AA041208 | 96 | subcomplex, 8 (19kD, ASHI) | #N/A | #N/A | 234.76 | 302.34 | 215.51 | 448.34 | 380 | 234.61 |
| rc_AA043790_at | AA043790 | 66 | KIAA0937 protein | #N/A | W/V# | 5.63 | 7.44 | 12.94 | 32.16 | 31.34 | 10.66 |
| AA044095_at | AA044095 | 102 | EST | #N/A | #N/A | 29.41 | 18.4 | 35.04 | 31.85 | 24.92 | 38.98 |
| | | | Autosomal Highly | | | | | | | | |
| AA044842_at | AA044842 | 105 | Conserved Protein | 5.21 | 0.000 | 16.66 | 16.52 | 18 | 167.15 | 159.32 | 123.77 |
| rc_AA045481_at | AA045481 | 107 | EST | #N/A | #N/A | 28.81 | 21.88 | 17.14 | 71.32 | 63.94 | 38.49 |
| rc AA046457 at | AA046457 | 11 | EST | 3.2 | ,0.00513 | 77.66 | 80.71 | 27.66 | 304.54 | 264.9 | 233.62 |
| rc_AA046747_at | AA046747 | 114 | EST | 4.82 | 0.00022 | -5.19 | 4.2 | 10.73 | 113.78 | 88.54 | 66.41 |
| AA047151_at | AA047151 | 116 | EST | 7.13 | 0.00007 | 17.55 | 17.5 | 10.09 | 188.62 | 185.41 | 80.5 |
| rc_AA053917_at | AA053917 | 131 | EST | #N/A | #N/A | -18.24 | -11.6 | 21.61 | 56.08 | 38.68 | 82.71 |
| rc_AA055992_at | AA055992 | 136 | calumenin | 3.51 | 0.00604 | 80.45 | 65.8 | 47.34 | 276.06 | 265.13 | 141.34 |
| AA056319_at | AA056319 | 139 | EST | #N/A | #N/A | 21.89 | 17.62 | 12.89 | 29.23 | 26.53 | 11.96 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | 704 | | | metastati metastati | metastati | normal | normal | norma |
|--|----------------------------------|-------------------|---|-----------------------|-------------------------|-------------------------|-------------------------|--------------------------|----------------------------|----------------------------|-------------------------|
| | | | | 2 | | , | וומנשפותנו | | | | |
| | | | | change | _ | metastati | ຮ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc_AA056482_at | AA056482 | 141 | EST | 4.82 | 0.00199 | 10.09 | 19.17 | 14.96 | 135.83 | 132.94 | 97.88 |
| rc_AA056735_at | AA056735 | 142 | SEC24 (S. cerevisiae) related gene family, member D | #N/A | Y / N # . | 4.59 | 10.03 | 14.42 | 29.84 | 28.43 | 10.97 |
| rc_AA074885_at rc_AA075298_at rc_AA076672_at | AA074885 AA075298 AA076672 | 161 163 172 | macrophage receptor with collagenous structure EST EST | 11.05 #N/A #N/A | 0.00786 #N/A #N/A | 79.55 46.45 86.18 | 25.58 36.93 87.53 | 153.64 42.61 60.88 | 652.03 129.13 122.82 | 761.74 121.57 105.84 | 300.57 70.36 54.8 |
| rc_AA084286_at rc_AA084318_at | AA084286 AA084318 AA086201 | 176 177 185 | paternally expressed gene 3 EST EST | #N/A #N/A 5.8 | #N/A #N/A 0.00012 | -2.06 9.73 21.29 | 0.32 3.97 24.75 | 14.88 17.06 14.37 | 30.7 30.63 177.39 | 26.84 32.37 182.95 | 20.69 11.48 95.51 |
| AA092376_at | AA092376 | 196 | 15 kDa selenoprotein hone mombogenetic profein | #N/A | #N/A | 16.78 | 14.2 | 28.1 | 59.07 | 57.94 | 24.97 |
| AA092596_at | AA092596 | 197 | 6 HLA-B associated transcript- | 3.46 | 0.02532 | 30.18 | 22.69 | 61.67 | 148.47 | 171.15 | 77.57 |
| AA092716 at | AA092716 | 198 | ဗ | 13.97 | 0.0000 | 62.83 | 63.53 | 42.33 | 952.09 | 817.41 | 545.31 |
| rc AA098864 at | AA098864 | 205 | EST | #N/A | #N/A | 30.42 | 31.06 | 18.38 | 56.43 | 52.57 | 21.3 |
| rc_AA099225_at | AA099225 | 206 | EST | 7.33 | 0.00062 | 4.37 | 1.35 | 90.9 | 212.68 | 163.45 | 194.31 |
| rc_AA099571_at | AA099571 | 209 | MD-2 protein | #N/A | #N/A | 10.12 | 4.61 | 19.11 | 55.77 | 62.72 | 31.71 |
| rc_AA102098_at | AA102098 | 218 | EST | #N/A | #N/A | 1 .8 | -5.18 | 15.21 | 21.79 | 20.01 | 6.78 |
| rc_AA102571_at | AA102571 | 220 | EST | #N/A | #N/Y | 8.7 | 12.32 | 15.12 | 17.15 | 15.33 | 7.6 |
| rc_AA112209_s_at AA112209 | AA112209 | 223 | acyl-Coenzyme A dehydrogenase, long chain | 3.37 | 0.00084 | 29.77 | 28.31 | 11.33 | 116 | 100.18 | 66.07 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|---------------------------|----------|--------|---|----------|---------|-----------|---------------------|-----------|--------|--------|---------|
| | | | | change | | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc AA115933 s at | AA115933 | 231 | KIAA1098 protein | W/W# | Y/N# | 8.62 | 6.46 | 10.77 | 33.88 | 34.38 | 16.63 |
| rc AA116075 at | AA116075 | 234 | EST | #N/A | W/V# | 29.71 | 23.88 | 10 | 62.94 | 64.79 | 25.13 |
| rc AA121140 at | AA121140 | 235 | EST | 3.33 | 0.00058 | 6.67 | 5.82 | 3.73 | 74.84 | 83.56 | 31.9 |
| rc_AA121257_at | AA121257 | 236 | EST | #N/A | #N/A | 21.76 | 21.21 | 32.49 | 42.39 | 43.62 | 11.55 |
| rc_AA126059_at | AA126059 | 246 | EST | 3.08 | 0.00706 | 102.34 | 104.33 | 40.8 | 380.92 | 280.71 | 316.4 |
| rc_AA127514_at | AA127514 | 253 | EST | 3.4 | 0.00045 | 11.36 | 10.76 | 8.9 | 74.71 | 76.5 | 36.07 |
| 1 | | | schwannomin interacting | | | | | | | | - |
| rc_AA127646_at | AA127646 | 254 | protein 1 | #N/A | #N/A | 9.79 | -0.26 | 12.36 | 44.24 | 41.48 | 20.02 |
| 1 | | | sequence-specific single- stranded-DNA-binding | | | | | | | | |
| rc AA128177 at | AA128177 | 258 | protein | #N/A | #N/A | 11.71 | 6.05 | 16.96 | 44.4 | 33.57 | 33.54 |
| rc_AA129465_f_at | AA129465 | 263 | EST | #N/A | #N/A | 7.9 | 10.16 | 24.29 | 61.62 | 59.75 | 34.37 |
| | | | meningioma expressed antigen 6 (coiled-coil proline- | | | | | | | | |
| rc AA133214 s at AA133214 | AA133214 | 276 | rich) | #N/A | #N/A | 3.16 | 7.46 | 14.35 | 65.17 | 56.11 | 51.18 |
| i I | | | calcitonin receptor-like receptor activity modifying | | | | | | | | |
| rc_AA133215_at | AA133215 | 277 | protein 1 | 4.55 | 0.02092 | 76.1 | 37.86 | 72.97 | 250.94 | 266.82 | 64.2 |
| rc AA133457 at | AA133457 | 280 | EST | #N/A | #N/A | 233.92 | 279.32 | 161.44 | 456.09 | 475.45 | 114 |
| rc_AA136333_at | AA136333 | 300 | zinc finger protein | #N/A | #N/A | 17.23 | 17.21 | 2.7 | 45.55 | 39.88 | 25.1 |
| rc_AA136611_at | AA136611 | 303 | EST | #N/A | #N/A | 9.05 | 6.82 | 8.39 | 48.23 | 42.67 | 13.53 |
| rc AA136940 at | AA136940 | 302 | EST | #N/A | #N/A | 40.81 | 41.78 | 20.7 | 79.13 | 73.37 | 41.89 |
| AA143019 at | AA143019 | 309 | EST | 6.75 | 0.00109 | -0.4 | -5.82 | 18.62 | 192.42 | 176.13 | 136.81 |
| rc_AA147626_at | AA147626 | 316 | EST | #N/A | #N/A | 37.93 | 20.43 | 44.09 | 82.38 | 75.98 | 37.45 |
| rc_aa147646_s_at AA147646 | AA147646 | 317 | DKFZP586A0522 protein flavin containing | 21.82 | 0 | 15.59 | 14.85 | 24.51 | 610.52 | 685.45 | 288.9 |
| rc_AA148480_s_at AA148480 | AA148480 | 318 | monooxygenase 5 | 19.64 | 0 | 18.26 | 14.26 | 14.85 | 521.95 | 407.11 | 247.99 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastat | metastati | normal | normal | normal |
|--------------------|----------|--------|---|--------------|----------|-----------|--------------------|---------------|--------|-----------------|---------|
| | | | | change | | metastati | ö | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | In metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc AA148539 at | AA148539 | 319 | EST | #N/A | #N/A | 96.6 | 9.87 | 8.65 | 28.88 | 25.89 | 6.73 |
| rc_AA149253_at | AA149253 | 323 | EST | 5.12 | 0.00863 | 78.65 | 75.3 | 61.14 | 401.77 | 301.06 | 333.03 |
| rc_AA150205_at | AA150205 | 328 | EST | #N/A | #N/A | 7.7 | -1.72 | 9.15 | 8.12 | 7.44 | 9.59 |
| rc_AA150284_at | AA150284 | 329 | EST | #N/A | #N/A | 28.51 | 25.35 | 17.97 | 41.67 | 43.94 | 18.29 |
| rc_AA151243_at | AA151243 | 334 | EST | #N/A | #N/A | 1.13 | -0.17 | 14.55 | 43.61 | 42.74 | 11.08 |
| | | | S-adenosylhomocysteine | | | | | | | | |
| AA157401 at | AA157401 | 346 | hydrolase-like 1 | #N/A | #N/A | 15.44 | 13.27 | 14.06 | 61.17 | 65.58 | 27.48 |
| rc AA167550 at | AA167550 | 361 | EST | #N/A | #N/A | 4.38 | 7.64 | 6.98 | 28.69 | 26.29 | 16.06 |
| rc_AA171529_at | AA171529 | 365 | EST | #N/A | #N/A | -1.76 | -4 .58 | 6.37 | 51.3 | 47.89 | 33.98 |
| AA174202 at | AA174202 | 375 | EST | #N/A | #N/A | 55.99 | 38.87 | 72.55 | 120.99 | 128.88 | 62.33 |
| rc AA179004 at | AA179004 | 377 | EST | 14.34 | 0.00008 | -33.2 | -28.97 | 78.62 | 503.76 | 495.87 | 326.16 |
| rc_AA182030_at | AA182030 | 387 | EST | 8.32 | 0.00018 | 14.05 | 16.82 | 12.13 | 222.23 | 220.01 | 117.56 |
| i I | | | STAT induced STAT | | | | | | | | |
| rc_AA182568_at | AA182568 | 388 | inhibitor-2 | 10.92 | 0.00099 | 23.5 | 18.15 | 21.15 | 501.87 | 386.2 | 478.02 |
| rc_AA187437_at | AA187437 | 389 | EST | #N/A | #N/A | 33.59 | 47.3 | 26.91 | 62.23 | 63 | 25.01 |
| | | | protein phosphatase 2 (formerly 2A), regulatory subunit A (PR 65), beta | | | | | | | | . |
| · rc_AA191310_s_at | | 397 | isoform | 7.28 #N/A | 0 4/1/4 | 26.23 | 29.87 | 6.93 23 54 | 206.65 | 209.15 38.29 | 77.34 |
| rc_AA193671_at | AA193071 | 403 | | ¥ | (| 5.0 | 1 | 0.00 | | 3.00 | 2 |
| AA195179 s at | AA195179 | 415 | eukaryotic translation initiation factor 4A. isoform 2 | W/W# | K/N# | 30.71 | 38.24 | 25.48 | 85.7 | 87.87 | 42.11 |
| rc AA195463 at | AA195463 | 416 | EST | | #N/A | 5.44 | 6.92 | 3.77 | 51.81 | 64.76 | 26.85 |
| rc_AA195515_at | AA195515 | 417 | EST | #N/A | #N/A | 7.27 | 3.56 | 8.61 | 51.59 | 49.91 | 23.13 |
| rc_AA195657_at | AA195657 | 419 | EST | 6.44 | 0.00016 | 5.72 | 7.44 | 9.74 | 157.7 | 136.21 | 100.68 |
| | | | | | | | | | | | |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affy ID | Genbank | Seq ID | Known Gene Name | fold change in metas | p value | metastati c: Mean | metastati metastati c: c: Std Median Dev | metastati c: Std Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|------------------------------|----------------------|------------|--|----------------------------|-----------------|----------------------|--|----------------------------|--------------------------|----------------------------|-----------------------------|
| . : | | ! | intercellular adhesion molecule 1 (CD54), human | | | | | | | 1 | |
| rc_AA197311_s_at | AA197311 | 422 | rhinovirus receptor | 6.07 | 0.00053 | -19.49 | 1.63 | 43.72 | 154.69 | 175.76 | 83.19 |
| rc_AA199603_at | AA199603 | 423 | EST | Y/V# | #N/A | 14.97 | 24.27 | 28.57 | 54.95 | 22.7 | 19.61 |
| rc_AA211370_at | AA211370 | 432 | EST | Y/V# | #N/A | 27.26 | 29.44 | 16.5 | 44.85 | 45.87 | 21.25 |
| rc AA211418 at | AA211418 | 434 | EST | #N/A | #N/A | 65.81 | 70.1 | 37.84 | 223.56 | 214.42 | 151.4 |
| rc_AA223902_at | AA223902 | 450 | EST | 9.91 | 0.00003 | 7.98 | -3.32 | 28.42 | 292.86 | 294.78 | 156.98 |
| rc_AA226925_at | AA226925 | 452 | EST | #N/A | #N/A | 11.94 | 9.22 | 7.55 | 49.69 | 48.29 | 27.49 |
| rc_AA227480_s_at | AA227480 | 456 | pim-2 oncogene | 3.31 | 0.02413 | 48 | 62.07 | 33.49 | 195.79 | 162.34 | 182.76 |
| rc_AA227968_at | AA227968 | 461 | histone deacetylase 6 epoxide hydrolase 2, | WW# | #N/A | 177.49 | 137.57 | 94.86 | 349.29 | 340.4 | 101.81 |
| rc AA232114 s at | AA232114 | 463 | cytoplasmic | 24.34 | 0.00007 | 63.09 | 46.66 | 60.38 | 1455.28 | 1626.11 | 664.36 |
| rc AA233126 at | AA233126 | 466 | EST | #N/A | #N/A | 48.86 | 59.64 | 33.59 | 88.56 | 74.17 | 35.35 |
| rc_AA233152_at | AA233152 | 467 | EST | 12.95 | 0 | -29.09 | -28.01 | 34.42 | 299.54 | 291.48 | 156.26 |
| AA233225_at | AA233225 | 468 | MRS1 protein | #N/A | #N/A | 25.16 | 30.66 | 26.03 | 54.1 | 55.8 | 22.69 |
| rc_AA233369 at | AA233369 | 471 | histidine ammonia-lyase | 90.6 | 0.0008 | 49.39 | 47.92 | 39.8 | 425.35 | 405.81 | 214.85 |
| rc_AA233763_at | , AA233763 | 472 | EST | 4.61 | 0.00004 | 25.29 | 36.26 | 17.49 | 146.52 | 139.66 | 60.53 |
| rc AA233797 at | AA233797 | 473 | sperm associated antigen 7 | #N/A | #N/A | 54.72 | 50.23 | 11.36 | 116.14 | 127.3 | 54.24 |
| rc_AA233837_at | AA233837 | 474 | EST | 4.79 | 0.0034 | 18.96 | 19.45 | 40.67 | 214.77 | 118.28 | 278.62 |
| | | , | CCAAT/enhancer binding | | 9 | 6 | 9 | 000 | 2.00 | 0 | 000 |
| AA234634_f_at AA234687_at | AA234634 AA234687 | 486 487 | protein (C/EBP), deita EST | 84.7 #N/A | 0.03518 #N/A | -8.8 -8.8 | 49.12 -12.38 | 20.12 | 61.87 | 286.94 47.79 | 532.04 |
| rc_AA234717_at | AA234717 | 489 | EST | #N/A | #N/A | 10.99 | 5.01 | 20.04 | 50.22 | 53.32 | 31.32 |
| AA234817_at | AA234817 | 490 | EST | 6.22 | 0.00099 | 31.51 | 20.97 | 34.92 | 222.41 | 156.99 | 133.06 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastat | metastati | normal | normal | normal |
|------------------|----------|--------|---|----------|---------|-----------|--------------------|-----------|--------|--------|---------|
| | | | | change | | metastati | ü | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Mrown Gene Name | in metas | | U | Median | Dev | Mean | Median | Std Dev |
| rc_AA234831_at | AA234831 | 491 | | 3.42 | 0.00206 | 23.54 | 21.62 | 28.81 | 112.19 | 119 | 49.23 |
| rc_AA235288_at | AA235288 | 494 | FIFLI-associated KnoGAF | 3.7 | 0.00643 | 34.06 | 30.95 | 14.36 | 169.9 | 113.81 | 138.05 |
| rc AA235507 at | AA235507 | 498 | golgi autoantigen, golgin suhfamily a. 5 | 3.28 | 0.00249 | 22.14 | 37.31 | 26.61 | 111.91 | 126.75 | 58.39 |
| rc_AA242822_at | AA242822 | 524 | EST | W/A | #N/A | 9.44 | 13.66 | 8.25 | 27.21 | 25.96 | 16.64 |
| rc_AA243654_at | AA243654 | 532 | EST | #N/A | #N/A | 0.67 | 3.93 | 11.08 | 69.19 | 62.85 | 48.63 |
| AA247453 at | AA247453 | 533 | EST | 3.09 | 0.0015 | 32.38 | 37.86 | 21.04 | 120.43 | 133.44 | 58.12 |
| rc_AA250958_f_at | AA250958 | 538 | EST | #N/A | #N/A | 53.41 | 53.63 | 22.55 | 99.74 | 114.71 | 55.23 |
| rc_AA251114_at | AA251114 | 539 | prostate cancer overexpressed gene 1 | 9.9 | 0.00039 | 28.47 | 18.55 | 28.89 | 219.81 | 202.99 | 87.55 |
| rc AA251776 at | AA251776 | 545 | jun D proto-oncogene | #N/A | #N/A | 30.8 | 23.59 | 32.23 | 51.25 | 46.83 | 15.96 |
| rc_AA251845_at | AA251845 | 548 | | #N/A | #N/A | 269.35 | 283.55 | 60.62 | 477.47 | 411.47 | 377.99 |
| rc_AA253410_at | AA253410 | 564 | EST | #N/A | #N/A | 18.46 | 7.23 | 42.79 | 49.37 | 26.82 | 45.2 |
| rc_AA255546_at | AA255546 | 569 | EST | 4 | 0.00301 | 61.04 | 67.88 | 31.64 | 260.34 | 224.12 | 142.84 |
| rc_AA255903_at | AA255903 | 573 | CD39-like 4 | 2.67 | 0.01687 | 72.5 | 39.23 | 108.92 | 383.56 | 374.1 | 211.92 |
| rc_AA256341_at | AA256341 | 578 | EST | 7.37 | 0.00091 | 17.81 | 2.34 | 28.59 | 280.57 | 324.08 | 170.98 |
| rc_AA256990_at | AA256990 | 585 | EST | #N/A | #N/A | 8.43 | 11.36 | 27.79 | 15.63 | 16.56 | 6.83 |
| rc AA257057 s at | AA257057 | 586 | EST | 8.11 | 0.00379 | 42.36 | 16.78 | 47.02 | 451.86 | 462.6 | 343.05 |
| rc_AA258158_at | AA258158 | 588 | EST | #N/A | #N/A | 7.91 | 2.43 | 17.94 | 44.84 | 34.32 | 35.63 |
| rc_AA258353_at | AA258353 | 593 | EST | 5.28 | 0.00193 | 71.76 | 84 | 37.77 | 347.7 | 363.14 | 106.2 |
| rc_AA259064_at | AA259064 | 602 | EST | 13.15 | 0.00001 | 15.32 | -1.37 | 28.67 | 401.93 | 394.99 | 178.23 |
| rc_AA278670_at | AA278670 | 616 | EST | #N/A | #N/A | 7.44 | 11.47 | 12.63 | 54.24 | 49.98 | 28.69 |
| rc_AA278824_at | AA278824 | 619 | EST | #N/A | W/V# | 26.85 | 31.59 | 11.72 | 83.48 | 91.25 | 30.24 |
| rc_AA278853_at | AA278853 | 621 | EST | #N/A | #N/A | 10.28 | 11.7 | 21.49 | 42.68 | 43.68 | 17.64 |
| rc_AA279158_i_at | AA279158 | 623 | EST | #N/A | #N/A | 20 | 46.97 | 19.41 | 87.9 | 72.44 | 38.36 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | 700 | | | motactati matactati | matactati | normal | normal | normal |
|------------------|----------|--------|---|----------|---------|-----------|---------------------|-----------|--------|--------|---------|
| | | | | 2 | | | merasian | metastan | | 3 | |
| | | | | change | | metastati | ö | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc AA279341 at | AA279341 | 625 | EST | #N/A | #N/A | 67.56 | 80.37 | 50.19 | 132.89 | 118.11 | 67.28 |
| rc_AA279916_at | AA279916 | 633 | EST | W/N# | #N/A | 61.08 | 56.18 | 25.7 | 115.24 | 102.89 | 58.77 |
| | • | | spleen focus forming virus (SFFV) proviral integration | | | | | | | | |
| rc_AA280413_s_at | AA280413 | 638 | oncogene spi1 | 4.46 | 0.02062 | 64.05 | 69.15 | 51.55 | 339.15 | 353.53 | 235.59 |
| rc_AA281545_at | AA281545 | 645 | EST | 3.64 | 0.00002 | 6.52 | -0.81 | 16.89 | 87.27 | 85.27 | 34.29 |
| | | | seven in absentia | | | | | | | | |
| rc_AA281770_at | AA281770 | 649 | (Drosophila) homolog 1 mannose-P-dolichol | 3.96 | 0.00094 | 7.67 | 4.72 | 15.41 | 103.75 | 79.4 | 70.74 |
| rc_AA281796_at | AA281796 | 650 | utilitzation defect 1 | 3.3 | 0.04108 | 65.08 | 53.58 | 57.86 | 170.88 | 165.02 | 41.87 |
| rc_AA282541 at | AA282541 | 661 | EST | #N/A | #N/A | 7.18 | 5.97 | 14.25 | 31.31 | 29.9 | 13.53 |
| rc_AA282956_at | AA282956 | 664 | EST | W/V# | #N/A | 0.28 | -9.07 | 30.81 | 59.89 | 51.54 | 35.5 |
| rc_AA283066_at | AA283066 | 999 | EST | #N/A | #N/A | 21.18 | 25.64 | 13.39 | 67.44 | 59.46 | 35.89 |
| | | | phosphatidylethanolamine N- | | | | | | | | |
| rc AA284795 at | AA284795 | 678 | methyltransferase | 10.03 | 0.00019 | 44.8 | 62.07 | 44.12 | 514.93 | 591.52 | 206.4 |
| rc_AA285053_at | AA285053 | 681 | EST | 6.95 | 0.00125 | 12.65 | 14.54 | 23.53 | 238.16 | 242.27 | 169.12 |
| rc_AA286710_at | AA286710 | 683 | lymphocyte adaptor protein | #N/A | #N/A | 37.88 | 39.15 | 30.32 | 82.93 | 86.15 | 45.58 |
| rc_AA287566_at | AA287566 | 069 | KIAA0187 gene product | 9.07 | 0.00013 | 4.86 | 6.24 | 7.4 | 246.24 | 201.66 | 228.64 |
| rc_AA291293_at · | AA291293 | 698 | EST | #N/¥ | #N/A | 17.35 | 18.07 | 17.24 | 33.6 | 33.3 | 5.56 |
| AA292440_s_at | AA292440 | 709 | | #N/A | #N/A | 122.1 | 125.27 | 37.73 | 370.32 | 318.31 | 181.37 |
| AA296821_at | AA296821 | 723 | EST | #N/A | #N/A | 15.47 | 10.65 | 21.91 | 57.72 | 71.28 | 33.61 |
| AA298180_at | AA298180 | 726 | EST | 3.1 | 0.00747 | 19.6 | 25.9 | 18.83 | 109.91 | 84.28 | 88.07 |
| rc_AA312946_s_at | AA312946 | 731 | EST | 9.21 | 0.00106 | 12.32 | 13.78 | 12.24 | 300.22 | 304.56 | 213.18 |
| | | ٠ | | | | | | | | | |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|----------------|------------------|------------|--------|--|----------|-----------|-----------|---------------------|-----------|---------|---------|---------|
| | | | | | change | - | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| | Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| | | | | | | | | | | | | |
| < | A A 246070 at | A A 346279 | 737 | ratty-acid-Coenzyme A licase Iong-chain 3 | ₩/N# | #W/A | 34.62 | 38.08 | 17.08 | 71.9 | 73.01 | 27.47 |
| { < | AA228684 at | AA328684 | | EST | #N/A | #N/A | 49.88 | 48.15 | 3141 | 115.97 | 114.66 | 46.25 |
| | F AA342918 at | AA342918 | | EST. | √N# | √× VN# | 23.67 | 31.68 | 25.77 | 69.4 | 73.71 | 18.13 |
| ر ک ا | | AA343142 | | EST | 20.87 | 0.00003 | -12.34 | -21.46 | 30.45 | 610.64 | 636.83 | 438.33 |
| | | | | 8 thenormon the molanda | | - | | | | | | |
| Δ. | r 003/1866 e at | A A 344866 | 752 | camma polypentide | 7.28 | 0.00206 | 292 65 | 305.28 | 176.61 | 1845.16 | 1679.29 | 561.04 |
| ל בי אל | rc_AA363203_s_at | AA363203 | 761 | EST | W.W.# | W/N# | 35.7 | 40.27 | 34.51 | 78.45 | 75.11 | 47.8 |
| . 4 | rc AA365691 at | AA365691 | 763 | EST | #N/A | #N/A | 48.01 | 47.17 | 25.37 | 28.69 | 26.73 | 13.89 |
| ב ל ב | rc_AA381125_at | AA381125 | 772 | EST | 15.48 | 0 | 17.66 | 13.82 | 13.2 | 412.26 | 344.45 | 217.56 |
| įΫ́ | AA397841 at | AA397841 | 780 | EST | 8.21 | 0 | 7.72 | 3.33 | 18.47 | 214.17 | 189.93 | 116.41 |
| rc A | rc AA397904 at | AA397904 | 781 | EST | #N/A | #N/A | 22.74 | 18.55 | 19.05 | 59.75 | 54.83 | 32.24 |
| ် မ မ မ | rc_AA397919_at | AA397919 | 785 | EST | #N/A | W/A# | 108.63 | 144.35 | 84.25 | 243.79 | 199.62 | 185.4 |
| | I | | | | | | | | | | | |
| | | | | growth factor receptor- | | | | | | | | |
| rc AA | rc AA398124 s at | AA398124 | 787 | bound protein 14 | 7.82 | 0.0000 | 3.4 | 5.86 | 7.61 | 189.27 | 167.23 | 110.44 |
| <u>ရ</u> | rc AA398280 at | AA398280 | 792 | EST | 12.43 | 0.00134 | -114.74 | -71.05 | 103.33 | 433.45 | 423.73 | 356.61 |
| ر ک | rc_AA398386_at | AA398386 | 793 | EST | 5.71 | 0.00007 | 10.59 | 16.25 | 21.59 | 153.16 | 164.38 | 83.94 |
| ် (၁ | rc_AA398423_at | AA398423 | 795 | EST | 8.26 | 0.00063 | -17.3 | -16.19 | 23.92 | 230.91 | 250.5 | 156.29 |
| ¹ ည | rc_AA398674_at | AA398674 | 798 | thrombospondin 1 | #N/A | #N/A | -27.87 | 14.49 | 146.65 | 137.87 | 96.14 | 119.21 |
| ۱ ک | rc_AA400030_at | AA400030 | 806 | EST | 3.98 | 0.00088 | 8.99 | 12.21 | 11.35 | 97.83 | 115.15 | 52.04 |
| ا 2 | rc_AA400080_at | AA400080 | 807 | EST | #N/A | W/A | 25.33 | 23.57 | 27.54 | 61.21 | 60.21 | 29.91 |
| Α Ι | rc_AA400258_at | AA400258 | 812 | EST | 11.89 | 0.00478 | 85.31 | 36.81 | 139.93 | 827.48 | 884.35 | 562.14 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affy ID | Genbank | Seq ID | Known Gene Name | fold change in metas | p value | metastati c: Mean | metastati metastati c: c: Std Median Dev | metastati c: Std Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|------------------|----------|--------|--|----------------------------|---------|----------------------|--|----------------------------|--------------------------|----------------------------|-----------------------------|
| | | | | - | | | | | | | |
| | | | tumor necrosis factor alpha- inducible cellular protein | | | | | | | | |
| | ٠ | • | containing leucine zipper domains; Huntingtin | | | | | | | | |
| | | | interacting protein L; transcrption factor IIIA- | | | | | ٠ | | | |
| AA400333_at | AA400333 | 815 | interacting protein | #N/A | #N/A | 27.22 | 27.33 | 24.37 | 62.54 | 58.6 | 31.69 |
| rc_AA400934_at | AA400934 | 824 | EST | 4.98 | 0.02013 | 75.44 | 67.21 | 7.77 | 305.09 | 307.45 | 125.76 |
| | | | calcitonin receptor-like receptor activity modifying | | | | | | | | |
| rc_AA400979_at | AA400979 | 825 | protein 3 | 6.65 | 0.01051 | 48.89 | 21.04 | 80.4 | 276.38 | 267.3 | 88.7 |
| rc_AA401091_at | AA401091 | 826 | EST | W/V# | #N/A | 31.88 | 32.59 | 43.51 | 30.55 | 19.82 | 28.68 |
| rc_AA401562_s_at | AA401562 | 830 | EST | 50.45 | 0.00301 | 155.46 | 63.87 | 317.6 | 3745.71 | 3628.4 | 1635.98 |
| rc_AA401825_at | AA401825 | 831 | EST | #N/A | #N/A | 24.3 | 26.82 | 18.04 | 65.22 | 57.51 | 46.54 |
| | • | | growth arrest and DNA- | | , | | | | | | |
| rc_AA402224_at | AA402224 | 836 | damage-inducible, gamma | 14.41 | 0.00012 | 37.55 | 48.88 | 48.58 | 749.36 | 812.2 | 443.66 |
| rc_AA402610_at | AA402610 | 839 | KIAA0548 protein | #N/A | #N/A | 3.02 | 69.9 | 13.83 | 27.54 | 23.2 | 17.78 |
| rc_AA402656_at | AA402656 | 841 | EST | 12.05 | 0.00001 | -2.56 | 18.8 | 40.57 | 342.11 | 315.47 | 205.61 |
| AA404214_at | AA404214 | 846 | EST | #N/A | #N/A | 35.28 | 48.54 | 23.52 | 99'02 | 66.71 | 55.97 |
| rc_AA404248_at | AA404248 | 847 | EST | #N/A | #N/A | 11.96 | 11.18 | 10.84 | 40.92 | 32.06 | 17.49 |
| rc_AA404352_at | AA404352 | 820 | EST | 7 | 0.00059 | 26.7 | 20.02 | 33.28 | 213.01 | 172.11 | 108 |
| rc_AA405494_at | AA405494 | 828 | EST | #N/A | #N/A | 10.03 | 9.53 | 7.62 | 59.95 | 44.88 | 50.8 |
| rc_AA405495_at | AA405495 | 829 | EST | #N/A | #N/A | 26.12 | 22.63 | 43.42 | 79 | 87.8 | 30.16 |
| rc_AA405744_at | AA405744 | 863 | EST | #N/A | #N/A | 61.18 | 28.74 | 58.71 | 64.39 | 57.95 | 35.67 |
| rc_AA406363_at | AA406363 | 874 | EST | #N/A | #N/A | 6.85 | 5.69 | 15.69 | 35.62 | 35.06 | 13.69 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|----------------|----------|--------|------------------------------|----------|---------|-----------|---------------------|-----------|--------|---------|---------|
| | | | | change | - | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | C Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc AA406546 at | AA406546 | 879 | EST | #N/A | #N/A | -1.45 | -7.89 | 15.19 | 46.62 | 50.65 | 26.97 |
| | AA406610 | 880 | EST | #N/A | #N/A | 5.99 | 6.04 | 4.63 | 35.94 | 34.84 | 11.57 |
| rc_AA410255_at | AA410255 | 882 | EST | 7.56 | 0.00043 | 0.82 | 4.15 | 11.49 | 195.75 | 236.93 | 110.49 |
| rc_AA411795_at | AA411795 | 892 | EST | #N/A | #N/A | 72.18 | 53.17 | 44.28 | 130.27 | 131.65 | 23.71 |
| rc_AA412063_at | AA412063 | 895 | | 8.26 | 0.00001 | 11.36 | 14.03 | 19.55 | 220.12 | 187.85 | 149.43 |
| rc_AA412068_at | AA412068 | 896 | | #N/A | #N/A | 30.52 | 29.79 | 13.4 | 81.26 | 74.52 | 35.68 |
| | | | | | | | | | | | |
| rc_AA412149_at | AA412149 | 897 | KIAA0480 gene product | #N/A | #N/A | 14.07 | 10.08 | 14.16 | 37.43 | 32.4 | 15.56 |
| rc_AA412520_at | AA412520 | 903 | EST | #N/A | W/V# | 18.84 | 14.4 | 14.11 | 80.29 | 103.19 | 41.98 |
| l I | | | ubiquitin-conjugating | | í | | | | | | |
| rc_AA412700_at | AA412700 | 904 | enzyme E2L 6 | #N/A | #N/A | 139.65 | 95.44 | 135.61 | 275.39 | 273.98 | 112.87 |
| | | | | | | | | | | | |
| | | | 5-methyltetrahydrofolate- | | | | | | | | |
| | | | homocysteine | | | | i | 0 | 77 | . 0 | 700 |
| rc_AA416936_at | AA416936 | | | | 0.00632 | 82.04 | 85.81 | 69.18 | 427.41 | 366.53 | 235.63 |
| rc_AA417078_at | AA417078 | | | 4.1 | 0.00414 | 36.75 | 35.35 | 15.12 | 189.88 | 171.1 | 150.02 |
| rc_AA418398_at | AA418398 | 921 | EST | #N/A | #N/A | 0.1 | -6.02 | 17.92 | 30.41 | 32 | 20.72 |
| rc_AA419608_at | AA419608 | 925 | EST | 9.19 | 0.00005 | 51.89 | 51.37 | 25.93 | 524.34 | 571.05 | 296.46 |
| rc_AA419622_at | AA419622 | 926 | EST | 4.62 | 0.00386 | 24.88 | 17.15 | 26 | 158.88 | 162.5 | 112.57 |
| rc_AA421051_at | AA421051 | 928 | serum-inducible kinase | #N/A | #N/A | 42.94 | 7.14 | 87.89 | 51.27 | . 45.65 | 28 |
| | | | hranched chain alpha- | | | | | | | | |
| | | | ketoacid dehydrogenase | | | | | | | | |
| rc_AA421052_at | AA421052 | 929 | kinase | 3.52 | 0.00869 | 77.76 | 94.75 | 43.63 | 251.45 | 221.35 | 109.93 |
| | | | insulin-like growth factor 2 | | | | | | | | |
| rc_AA421561_at | AA421561 | 933 | | 9.98 | 0.00007 | 79.46 | 77.34 | 41.63 | 921.91 | 703.16 | 679.72 |
| | | | | | | | | | | | |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|--|--|------------------------------|---|-------------------------------------|--|---------------------------------|--|--|---|--|---|
| | | | | change | | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| AA424307 at | AA424307 | 944 | EST | 5.73 | 0.0074 | 44.98 | 21.05 | 54.01 | 202.82 | 194.92 | 68.07 |
| rc AA424798 at | AA424798 | 947 | EST | 17.45 | 0.00352 | 48.63 | 15.48 | 171.26 | 879.91 | 873.25 | 489.59 |
| rc_AA425214_at | AA425214 | 950 | EST | #N/A | #N/A | 2.71 | 2.07 | 4.88 | 21.4 | 20.31 | 10.15 |
| rc_AA426643_at | AA426643 | 970 | EST | #N/A | #N/A | 9.96 | 0.97 | 18.11 | 48.26 | 47.37 | 23.47 |
| rc_AA427537_at | AA427537 | 974 | DKFZP566J153 protein regulator of G-protein | #N/A | #N/A | 183.26 | 156.56 | 68.58 | 259.57 | 242.97 | 79.81 |
| rc_AA427579_at | AA427579 | 975 | signalling 14 | #N/A | #N/A | 13.35 | 13.28 | 16.49 | 34.79 | 25.25 | 25.11 |
| rc_AA427819_at | AA427819 | 980 | midline 2 | 3.44 | 0.00063 | 25.98 | 17.06 | 14.91 | 97.03 | 93.49 | 31.39 |
| rc_AA428150_at | AA428150 | 985 | EST | 5.24 | 0.00167 | 41.92 | 35.06 | 30.53 | 213.96 | 217.27 | 76.08 |
| rc_AA428325_at | AA428325 | 988 | EST | 8.36 | 0.00002 | -0.52 | 3.28 | 16.92 | 194.02 | 167.37 | 111.11 |
| rc_AA428900_at | AA428900 | 992 | EST | 7.01 | 0.00037 | 66.25 | 22.99 | 15.35 | 615.96 | 619.42 | 441.72 |
| rc_AA429038_at | AA429038 | 995 | EST | 3.29 | 0.00927 | 13.39 | 1.93 | 28.01 | 108.66 | 87.59 | 86.11 |
| rc AA429478 at | AA429478 | 866 | EST | 3.41 | 0.02599 | 55.86 | 65.19 | 47.48 | 192.7 | 196 | 89.2 |
| rc_AA429651_at | AA429651 | 1002 | KIAA0871 protein | #N/A | #N/A | 15.52 | 13.25 | 17.03 | 44.75 | 41.91 | 28.67 |
| rc_AA432166_f_at_AA432166 | AA432166 | 1030 | succinate dehydrogenase complex, subunit C, integral membrane protein, 15kD | #N/A | .) | 10.94 | 1.41 | 42.99 | 33.86 | 42.75 | 27.46 |
| rc_AA434225_at | AA434225 | 1035 seru | serum constituent protein | #N/A | #N/A | 1010.84 | 876.42 | 674.94 | 1022.5 | 746.49 | 681.49 |
| rc_AA435591_at rc_AA435753_at rc_AA436156_s_at rc_AA436548_at rc_AA436880_at | AA435591 AA435753 AA436156 AA436548 AA436880 | 1038 1045 1051 1054 | kinesin family member 3B EST EST EST EST | 3.5 4.71 #N/A #N/A 3.22 | 0.0001 0.00078 #N/A #N/A 0.00699 | 3.57 198.97 22.66 5.08 | 3.68 193.33 7.78 5.93 9.21 | 10.34 110.58 34.68 5.2 13.85 | 75.9 970.79 54.98 40.27 86.89 | 81.12 830.39 48.35 38.99 68.41 | 29.32 567.79 38.09 22.22 66.5 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affy ID | Genbank | Seq ID | Known Gene Name | fold change in metas | p value | metastati c: Mean | metastati metastati c: c: Std Median Dev | metastati c: Std Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|--|--|---|--|-------------------------------|--------------------------------------|-----------------------------------|--|---------------------------------|-------------------------------------|------------------------------------|------------------------------------|
| rc_AA437295_at | AA437295 | 1062 rit | 1062 ribosomal protein L7a | 4.35 | 0.00347 | 19.67 | 20.38 | 43.07 | 139.16 | 140.38 | 70.9 |
| rc_AA443658_at | AA443658 | tra 1079 su | transmembrane 7 superfamily member 2 | 9.06 | 0.00048 | 4.48 | 17.91 | 24.51 | 276.02 | 194.73 | 227.48 |
| rc_AA443934_at | AA443934 | 1083 G | GTP-binding protein Rho7 | 3.09 | 0.00214 | 30.32 | 35.97 | 13.76 | 115.87 | 113.7 | 82.99 |
| rc_AA446342_at rc_AA447802_at AA447876_at | AA446342 AA447802 AA447876 | sc 1088 (T 1108 E 1109 E | seven in absentia (Drosophila) homolog 1 EST EST | 4.84 #N/A #N/A | 0.00015 #N/A #N/A | 9.92 15.64 13.05 | 9.12 17.7 6.78 | 4.58 13.81 34.07 | 110.73 21.51 27.08 | 115.89 21.34 22.86 | 53.13 7.37 22.46 |
| rc_AA448300_at rc_AA449108_at rc_AA449297_at | AA448300 AA449108 AA449297 | FXYI trans 1116 (phos 1118 EST 1121 EST | FXYD domain-containing ion transport regulator 1 (phospholemman) EST | 24.97 #N/A 3.78 | 0.00001 #N/A 0.00039 | 118.64 10.9 3.76 | 81.63 18.13 0.63 | 70.34 15.28 19.01 | 2849.54 52.58 91.67 | 2905.51 57.3 81.15 | 994.41 23.76 55.17 |
| rc_AA452158_at rc_AA453770_s_at rc_AA454177_i_at rc_AA454667_at | AA452158 AA453770 AA454177 AA454667 | 1141 m 1157 E 1164 E 1167 E | ras homolog gene family, member B EST EST EST | 28.96 6.04 10.3 #N/A | 0.00064 0.00524 0.0008 #N/A | -103.87 46.33 9.16 21.25 | -106.99 25.95 14.72 12.77 | 71.68 53.45 11.39 23.4 | 1071.9 217.46 324.16 83.02 | 1164.8 212.4 253.23 75.81 | 789.64 70.02 258.82 56.63 |
| rc_AA455111_at | AA455111 | h 1173 ri | heterogeneous nuclear ribonucleoprotein C (C1/C2) | #N/A | #N/A | -31.71 | -40.39 | 38.45 | 38.96 | 36.83 | 26.21 |
| rc_AA455261_at | AA455261 | 1175 c | 1175 chromobox homolog 7 | #N/A | #N/A | 32.5 | 45.54 | 26.51 | 82.99 | 80.99 | 30.15 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| rc_AA455865_at AA455865 1180 class B rc_AA455986_s_at AA455896 1181 glypican 1 rc_AA455987_at AA455987 1183 EST rc_AA455988_at AA455988 1184 hydroxylass rc_AA456075_at AA456075 1186 homolog A rc_AA456080_at AA456080 1187 EST rc_AA456147_at AA456147 1188 IIIA rc_AA456812_at AA456612 1197 EST rc_AA456812_at AA456687 1197 EST rc_AA456823 AA456687 1197 EST rc_AA456852_at AA456845 1198 KIAA0680 rc_AA458652_at AA458652 1202 EST rc_AA458651_at AA458652 1202 EST rc_AA458652_at AA458652 1202 EST | Known Gene Name | change in metas | p value | metastati c: Mean | c: Median | c: c: Std Median Dev | set 2: Mean | set 2: Median | set 2: Std Dev |
|--|---|--------------------|---------|----------------------|--------------|-------------------------|-----------------|------------------|-------------------|
| AA455987 1183 AA455987 1183 AA456075 1186 AA456080 1187 AA456147 1189 AA456289 1189 AA456612 1195 AA456687 1197 AA456687 1197 AA458652 1202 AA458652 1202 AA459256 1210 AA459256 1212 | phosphatidylinositol glycan, class B | 5.41 | 0.00004 | 11.68 | 8.67 | 10.65 | 133.15 | 134.74 | 63.3 |
| AA455987 1183 AA456086 1187 AA456080 1187 AA456289 1189 AA456612 1195 AA456687 1195 AA456687 1197 AA458652 1202 | glypican 1 | 3.46 | 0.00887 | 16.1 | 4.35 | 40.39 | 120.52 | 137.84 | 66.93 |
| AA456075 1186 AA456147 1188 AA456289 1189 AA456612 1195 AA456687 1197 AA456687 1198 AA456845 1198 AA456852 1202 AA456651 1202 AA456651 1202 AA456651 1202 | EST | 5.36 | 0.00029 | 20.54 | 17.51 | 15.24 | 128.55 | 130.64 | 17.07 |
| AA456075 1186 AA456080 1187 AA456289 1189 AA456326 1191 AA456612 1195 AA456687 1195 AA456845 1198 AA456845 1202 AA459256 1210 AA459256 1210 | butyrobetaine (gamma), 2- oxoglutarate dioxygenase (gamma-butyrobetaine hydroxylase) | 15.54 | 0.00001 | 10.13 | 11.88 | 8:38 | 354.42 | 375.91 | 141.08 |
| AA456080 1187 AA456289 1189 AA456289 1189 AA456612 1195 AA456687 1197 AA456845 1198 AA456852 1202 AA459005 1210 AA459256 1212 | RAD23 (S. cerevisiae) | | 771477 | 70 | | c c | 7 | ç | 6 |
| AA456289 1189 AA456289 1189 AA456612 1195 AA456687 1197 AA456845 1198 AA456845 1202 AA459005 1210 AA459256 1212 AA459256 1212 | TS: | ₹XX# | ₹ 4/N# | 32.35 | 17.59 | 33.21 | 30.47 27.76 | 26.92 | 12.38 |
| AA456289 1189 AA456289 1189 AA456612 1195 AA456687 1195 AA456845 1198 AA458652 1202 AA458652 1202 AA459005 1210 AA459256 1212 | general transcription factor | | | | | | | | |
| AA456289 1189 AA456612 1195 AA456687 1195 AA456845 1198 AA458652 1202 AA458652 1202 AA459005 1210 AA459256 1212 | IIA | 4.23 | 0.00088 | 4.61 | 0.17 | 9.08 | 102.1 | 89.95 | 63.84 |
| AA456612 1195 AA456612 1195 AA456687 1197 AA456845 1198 AA458652 1202 AA459005 1210 AA459256 1212 AA460661 1229 | EST | 15.31 | 0.00004 | 18.07 | 17.84 | 39.22 | 512.64 | 542.52 | 303.15 |
| AA456612 1195 AA456687 1197 AA456845 1198 AA458652 1202 AA459005 1210 AA459256 1212 AA460661 1229 | EST | 3.35 | 0.00489 | 17.08 | 2.74 | 30.38 | 111.5 | 102.17 | 68.09 |
| AA456687 1197 AA456845 1198 AA458652 1202 AA459005 1210 AA459256 1212 AA460661 1229 | EST | #N/A | W/A | 164.11 | 136.36 | 117.01 | 254.26 | 266.45 | 84.11 |
| AA456845 1198 AA458652 1202 AA459005 1210 AA459256 1212 AA460661 1229 | EST | 3.08 | 0.01189 | 17.88 | 37.03 | 53.32 | 130.65 | 118.55 | 65.44 |
| AA459005 1210 AA459256 1212 AA460661 1229 | KIAA0680 gene product | #N/A | #N/A | .23.95 | 17.11 | 17.99 | 67.15 | 63.46 | 49.27 |
| AA459005 1210 AA459256 1212 AA460661 1229 | EST | 8.26 | 0.00001 | 19.3 | 18.21 | 17.18 | 203.23 | 218.67 | 53.52 |
| AA459256 1212 AA460661 1229 | EST | #N/A | #N/A | -3.57 | 12.22 | 40.05 | 49.76 | 42.58 | 28.85 |
| | lectin, mannose-binding, 1 EST | 3.01 | 0.00094 | 3.83 | 8.1 -6.03 | 8.32 | 65.91 184.62 | 63.36 | 39.92 |
| rc_AA461448_at AA461448 1240 EST | EST | #N/A | #N/A | 1.74 | 4.19 | 22.52 | 58.65 | 56.06 | 32.42 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|------------------|----------------------|--------|---|--------------|-----------------|----------------|---------------------|---------------|-----------------|-----------------|----------------|
| | | | | change | | metastati | ដ | c: Std | set Z: | set 2: | set Z: |
| Affv ID | Genbank | Sed ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc AA463729 at | AA463729 | 1250 | EST | 4.07 | 0.00676 | 19.31 | 19.05 | 13.37 | 150.62 | 116.86 | 156.67 |
| rc AA463876 at | AA463876 | | EST | 3.31 | 0.00109 | 9.81 | 11.89 | 10.39 | 73.95 | 73.76 | 46.64 |
| rc AA464606 at | AAA64606 | | MRS1 protein | #N/A | #N/A | 24.35 | 19.03 | 39.62 | 59.55 | 36.75 | 57.76 |
| rc_AA465381_at | AA465381 | | | #N/A | #N/A | 88.83 | 36.24 | 130.22 | 102.73 | 113.07 | 52.43 |
| rc_AA465720_at | AA465720 | | EST | #N/A | #N/A | 11.5 | 21.33 | 37.22 | 93.06 | 108.97 | 43.26 |
| rc_AA470153_at | AA470153 | 1275 | solute carrier family 21 (organic anion transporter), member 9 | 13.26 | 0.00315 | 47.49 | 48.57 | 116.54 | 726.75 | 713.23 | 297.62 |
| | | | homolog of mouse quaking QKI (KH domain RNA | ; | ; | ! | | , | . (| į | i C |
| rc_AA478104_at | AA478104 AA478441 | 1296 | binding protein) cathensin F | #N/A 5.07 | #N/A 0.00752 | -7.07 53.85 | -11.44 67.48 | 23.2 43.52 | 62.06 243.44 | 47.17 243.61 | 56.07 69.17 |
| rc_AA479148_at | AA479148 | | EST | 38.05 | 0 | 6.2 | 1.32 | 14.38 | 895.91 | 847.72 | 362.53 |
| AA479266_at | AA479266 | | EST | #N/A | #N/A | 18.04 | 21.7 | 25.73 | 42.53 | 38.32 | 18.46 |
| rc AA479488 at | AA479488 | 1313 | S-adenosylhomocysteine hydrolase-like 1 | 4 | 0.0269 | 75.18 | 60.93 | 59.8 | 241.1 | 222.96 | 112.87 |
| rc_AA479961_at | AA479961 | 1320 | EST | #N/A | #N/A | 33.49 | 28.45 | 10.58 | 88.13 | 89.35 | 36.9 |
| rc AA479968 s at | | 1321 | arylsulfatase A | 9.01 | 0.00224 | 37.97 | 20.36 | 45.8 | 331.32 | 312.63 | 97.49 |
| rc_AA480991_s_at | | 1323 | ı EST | 8.59 | 0.00156 | 48.08 | 25.74 | 51.03 | 444.29 | 309.38 | 367.89 |
| rc_AA480997_i_at | • | 1324 | MADS box transcription enhancer factor 2, polypeptide A (myocyte enhancer factor 2A) | YN# | YN# | 91 | 20.58 | 9.93 | 68.53 | 55.62 | 63.6 |
| rc_AA481057_f_at | AA481057 | 1325 | EST | W/N# | ¥N/¥ | 16.3 | 18.55 | χ. Γ. | 33.83 | 37.89 | 73.27 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|----------------------------------|----------------------|--------------|------------------------------|--------------|--------------|-----------|---------------------|-----------|-----------------|--------------|-----------------|
| | | | | change | | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank Seg ID | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc AA486407 at | AA486407 | 1347 | EST | #N/A | #N/A | 33.22 | 47.88 | 28.61 | 81.72 | 97.02 | 55.07 |
| rc_AA486567_at | AA486567 | 1350 | EST | വ | 0.00002 | 4.65 | 2.5 | 21.62 | 131.53 | 98.76 | 95.59 |
| rc_AA486794_at | AA486794 | 1351 | EST | #N/A | #N/A | 49.57 | 52.25 | 18.73 | 78.8 | 72.55 | 31.84 |
| rc_AA487195_at | AA487195 | 1354 | EST | W/V# | #N/A | 8.73 | 7.94 | 9.21 | 21.04 | 14.67 | 16.52 |
| rc_AA487503_at | AA487503 | 1356 | EST | 8.85 | 0.00012 | 13.8 | 17.99 | 8.76 | 220.45 | 255.47 | 119.11 |
| rc_AA487576_at | AA487576 | 1357 | EST | #N/A | #N/A | -12.17 | -14.87 | 17.17 | 26.05 | 25.27 | 11.13 |
| rc_AA489009_at | AA489009 | 1366 | EST | #N/A | #N/A | 15.19 | 18.01 | 8.67 | 59.19 | 45.15 | 41.42 |
| rc_AA490882_s_at | AA490882 | 1381 | EST | 3.29 | 0.00319 | 20.67 | 13.84 | 27.06 | 100.67 | 98.47 | 54.46 |
| rc_AA490890_at | AA490890 | 1382 | EST | 3.02 | 0.00007 | 20.26 | 21.34 | 9.23 | 75.61 | 65.02 | 33.37 |
| rc_AA490947_at | AA490947 | 1383 | EST | #N/A | #N/A | 34.15 | 35 | 21.49 | 56.24 | 38.75 | 49.85 |
| rc_AA490964_at | AA490964 | 1384 | EST | #N/A | #N/A | 20.3 | 17.53 | 11.39 | 47.7 | 41.84 | 25.18 |
| rc_AA495803_at | AA495803 | 1392 | EST | #N/A | #N/A | 95.29 | 83.46 | 90.72 | 94.07 | 93.85 | 37.65 |
| 1 10000 | 4 4 4 4 4 4 | , , | 00 | . 4/14/7 | 471477 | | c | Ċ | 00 | . 77 | 50 |
| rc_AA495924_at | AA495924 | 1380 | Kinesin lamily member 35 | ¥121# | 421 | 4.10 | 3.20 | 2.93 | 22.03 | 04.40 | 6.85 |
| rc_AA496053_at | AA496053 | 1396 | EST | 3.28 | 0.00095 | 5.36 | 2.22 | 17.58 | 81.79 | 94.01 | 42.77 |
| rc_AA496927_at | AA496927 | 1402 | EST | #N/A | #N/A | 24.89 | 26.85 | 9.47 | 58.2 | 58.42 | 17.68 |
| rc AA496936 at | AA496936 | 1403 | EST | W/W# | #N/A | 17.71 | 9.86 | 20.19 | 30.24 | 34.4 | 14.77 |
| rc_AA504324_at | AA504324 | 1412 | EST | #N/A | #N/A | 45.99 | 38.81 | 31.27 | 74.31 | 72.05 | 31.59 |
| rc_AA521290_at | AA521290 | 1421 | EST | 4.53 | 0.0148 | 46.54 | 22.84 | 63.63 | 166.84 | 174.65 | 58.28 |
| rc_AA598412_at | AA598412 | 1425 | EST | #N/A | #N/A | -2.4 | -5.93 | 29.83 | 23 | 52.53 | 29.41 |
| rc_AA598453_s_at | AA598453 | 1429 | EST | #N/A | #N/A | 7.27 | 7.06 | 9.67 | 67.15 | 54.25 | 41.65 |
| · | | | lectin, galactoside-binding, | | | | - | | | | |
| rc_AA598685_at | AA598685 | 1435 | soluble, 8 (galectin 8) | #N/A | #N/A | 16.45 | 11.35 | 11.87 | 55.9 | 55.08 | 42.18 |
| rc_AA599107_at | AA599107 | 1443 | EST | #N/# | #N/¥ | 53.34 | 6 | 88.85 | 77.28 | 58.08 | 40.75 |
| | | | endothelin converting | | | | | | | | |
| rc_AA599199_at rc_AA599214_at | AA599199 AA599214 | 1444 1446 | enzyme 1 FST | #N/# #N/# | 4/N# #N/A | 187.77 | 108.97 7.36 | 225.55 | 695.29 34.58 | 680.26 36 | 375.12 14.54 |
| 17.00 | 70000 | | | | | | ? | į |) | 3 | |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati | metastati metastati | normal | normal | normal |
|----------------------------------|----------------------|-----------------------------|--|--------------|-----------------|----------------|----------------|---------------------|-----------------|----------------|-----------------|
| | | | | change | _ | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affv ID | Genbank | Sed ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc AA599365 at | AA599365 | 1449 | 449 decorin | #N/A | #N/A | 36.41 | 34.43 | 21.53 | 83.84 | 84.81 | 43.41 |
| rc_AA600248_at | AA600248 | 1461 | EST | #N/A | #N/A | 59.11 | 59.91 | 32.47 | 80.55 | 71.14 | 61.42 |
| rc AA608546 at | AA608546 | | EST | 12.52 | 0.00003 | -19.59 | -29.07 | 29.39 | 310.07 | 300.44 | 189.6 |
| rc_AA608723_at | AA608723 | | EST | #N/A | #N/A | 17.44 | . 15.29 | 13.31 | 66.54 | 70.38 | 20.94 |
| | 070000 | , | | 1 | 7 | 9 00 | 74 64 | . E | 736 54 | 226 70 | 65.0 |
| rc_AA609316_at | AA609316 | | EGF-like-domain, multiple 5 | 9.9 | 0.0001 | 0.77 | 10.17 | 23.30 | 264 62 | 242.13 | 105.33 |
| rc_AA609519_at rc_AA609715_at | AA609519 AA609715 | 1482 1488 | EST EST | 8.13 #N/A | 0.00003 #N/A | -1.49 | -3.92 | 10.63 | 29.09 | 33.14 | 17.26 |
| rc_AA620965_at | AA620965 | 1511 | NOT3 (negative regulator of transcription 3, yeast) | #N/A | #N/A | 4.34 | 8.96 | 16.51 | 48.05 | 40.77 | 32.06 |
| | | | similar to Caenorhabditis | | | | | | | | |
| rc_AA621209_at | AA621209 | 1516 | elegans protein C42C1.9 | 6.34 | 0.00144 | 22.77 | 19.95 24.68 | 46.78 | 214.61 | 167.05 | 138.24 65.35 |
| rc_AAbz1z35_ar | AA621233 | 161 | EST catenin (cadherin- associated protein) alpha- | † † | 0.002 | 67.02 | 7.00 | |) - | 2 | |
| rc_AA621315_at | AA621315 | 1521 | like 1 | #N/A | #N/A | 191.28 | 141.04 | 182.69 | 313.23 | 281.93 | 134.57 |
| rc_AA621796_at AB000114_at | AA621796 AB000114 | 1531 1532 | 1531 kinesin family member 3B 1532 osteomodulin | 4.44 #N/A | 0.00032 #N/A | 21.81 -2.62 | 23.64 -0.81 | 8.64 29.14 | 128.01 31.94 | 124.81 25.4 | 70.04 20.87 |
| AF000573_ma1_at AF000573 | AF000573 | homc dioxy 1543 oxida | homogentisate 1,2- dioxygenase (homogentisate oxidase) | 13.76 | 0.00002 | 9.05 | 13.23 | 17.33 | 380.3 | 348.9 | 256.4 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| ! | | 2 | 2 | fold change | | metastati | metastati c: | metastati metastati c: c: Std | normal set 2: | normal set 2: | set 2: |
|--------------|----------------|--------------|--|----------------|---------|-----------|-----------------|----------------------------------|------------------|------------------|---------|
| Affy ID | Genbank Seq ID | Sed ID | Known Gene Name | in metas | p value | C: Mean | Median | 200 | | Median | ota Dev |
| | | | solute carrier family 4, sodium bicarbonate | | | | | | | | |
| AF007216_at | AF007216 | 1550 | cotransporter, member 4 | 5.79 | 0.00005 | 14.85 | 9.84 | 13.93 | 157.72 | 162.44 | 99.26 |
| C02532_at | C02532 | 1563 | ST | #N/A | Y/V# | 10.12 | 16.41 | 16.79 | 32.27 | 34.25 | 11.02 |
| C15871 at | C15871 | 1575 | ST | 3.26 | 0.00046 | 13.83 | 18.19 | 16.99 | 79.77 | 83.41 | 40.17 |
| C16420_s_at | C16420 | 1576 | ST | 5.92 | 0.00119 | 26.92 | 20.84 | 24.82 | 205.53 | 234.17 | 120.92 |
| l I | | | umor susceptibility gene | | | | | | | | |
| C18029_at | C18029 | 1577 1 | 01 | #N/A | #N/A | 27.46 | 26.72 | 26.09 | 94.81 | 95.79 | 33.89 |
| rc_C20653_at | C20653 | 1578 E | :ST | 10.59 | 0.00001 | 7.99 | 2.12 | 11.85 | 251.82 | 299.09 | 106.09 |
| | | | | | | | | | | | |
| | | ω <i>τ</i> π | aldehyde dehydrogenase 5 family member A1 | | | | | | | | |
| | | ت : | succinate-semialdehyde | | | | | | | | |
| rc_C20982_at | C20982 | 1582 d | dehydrogenase) | #N/A | #N/A | 68.8 | 85.38 | 28.73 | 104.44 | 88.12 | 66.28 |
| rc_C21130_at | C21130 | 1583 E | EST | 8.79 | 0.00008 | 17.56 | 8.6 | 19.19 | 277.18 | 238.8 | 188.55 |
| rc_D11802_at | D11802 | 1597 a | ıngiotensinogen | 5.65 | 0.00000 | 55.88 | 52.43 | 25.14 | 319.73 | 291.02 | 132.07 |
| | | | | | | | | | | | |
| | | 7 | low density lipoprotein | | | | | | | | |
| | | , | receptor (familial | 3 | 0000 | 1 | 6 | | 1 | 9 | |
| rc_D11835_at | D11835 | 1598 | 1598 hypercholesterolemia) | 21.76 | 0.00307 | 76.38 | 23.21 | 131.25 | 7.088 | 910.39 | 322.98 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affy ID | Genbank | Seq ID | Known Gene Name | fold change in metas | p value | metastati c: Mean | metastati metastati c: c: Std Median Dev | metastati c: Std Dev | set 2: Mean | set 2: Median | set 2: Std Dev |
|----------------------------|------------------|--------|---|----------------------------|---------|----------------------|--|----------------------------|------------------|------------------|-------------------|
| D12485_at | D12485 | 1600 | phosphodiesterase //nucleotide pyrophosphatase 1 (homologous to mouse Ly- 41 antigen) | 4.57 | 0.00008 | -2.69 | -0.53 | 9.86 | 101.7 | 90.07 | 46.62 |
| D12620 s_at | D12620 | 1601 | cytochrome P450, subfamily IVF, polypeptide 2, cytochrome P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) | 35.09 | 0.00015 | 41.08 | 36.71 | 10.11 | 604.7 | 631.98 | 249.32 |
| D13243_s_at | D13243 | 1602 | pyruvate kinase, liver and RBC | 20.22 | 0 | -35.85 | -34.85 | 51.7 | 579.28 | 445.53 | 502 |
| D13814_s_at D14012_s_at | D13814 D14012 | 1611 | angiotensin receptor 1,angiotensin receptor 1B HGF activator | 3.12 12.75 | 0.00101 | 13.86 160.15 | 10.01 216.05 | 12.82 107.03 | 79.69 1705.96 | 75.71 1963.23 | 45.03 794.42 |
| D14664_at | D14664 | 1616 | KIAA0022 gene product | 8.98 | 0.00011 | 14.15 | 13.51 | 9.07 | 233.08 | 248.97 | 111.44 |
| D14686_at | D14686 | 1617 | aminomethyltransferase (glycine cleavage system protein T) | #N/A | #N/A | 76.93 | 90.12 | 29.68 | 171.07 | 190.08 | 34.32 |
| D14695_at | D14695 | 1618 | KIAA0025 gene product; 1618 MMS-inducible gene | 6.48 | 0 | 28.53 | 28.21 | 12 | 196.04 | 197.12 | 77.25 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | fold change | | metastati | metastati metastati c: c: Std | metastati c: Std | normal set 2: | normal set 2: | normal set 2: |
|---|---|-------------------------------|--|--|-----------------------------------|------------------------------|-------------------------------------|-------------------------------------|-----------------------------------|
| Seq ID | Known Gene Name | ame in metas | tas p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| SA (| SA (rat hypertension- 1620 associated) homolog | 3.83 | 3 0.00117 | 7 27.34 | 29.05 | 12.27 | 128.95 | 133.42 | 74.27 |
| 1622 histic | 1622 histidine ammonia-lyase | ase 22.66 | 0 99 | 11.74 | 8.13 | 16.04 | 538.78 | 535.2 | 166.14 |
| 1622 histid | 1622 histidine ammonia-lyase | ase 22.66 | 0 99 | 12.06 | 22.91 | 55.95 | 289.6 | 256.22 | 113.59 |
| basic 1647 bindir | basic transcription element binding protein 1 | ement 5.35 | 5 0.00086 | 3 16.08 | 23.65 | 17.64 | 166.06 | 203.79 | 87.22 |
| reguci 1648 marke 1661 EST 1662 EST 1664 EST | ıcalcin (senescence ker protein-30) | 10.55 #N/A 3.82 5.64 | 55 0.00037 A #N/A 2 0.00193 4 0.00384 | 7 11.48 -19.35 3 12.24 4 34.3 | 10.25 -33.07 -3.36 29.86 | 7 27.77 29.18 13.09 | 298.48 31.63 126.63 301.07 | 281.03 32.56 109.06 218.14 | 183.56 42.02 70.64 282.7 |
| 1668 hyalur | 1668 hyaluronan-binding protein 2 | rotein 2 18.13 | 3 0.00012 | 2 11.38 | -0.72 | 41.45 | 509.66 | 531.78 | 127.5 |
| potass rectify 1669 J, mer | potassium inwardly- rectifying channel, subfamily J, member 8 | ubfamily #N/A | A #N/A | -2.39 | 3.94 | 24.71 | 76.55 | 72.56 | 48.96 |
| potassium inw rectifying chan 1670 J, member 11 1687 EST | potassium inwardly- rectifying channel, subfamily J, member 11 EST | ubfamily #N/A #N/A | A #N/A A #N/A | 61.44 | 65.47 6.61 | 28.34 9.79 | 70.38 77.62 | 76.35 55.56 | 25.4 72.14 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| normal normal | set 2: set 2: | Median Std Dev | | | 103.85 69.38 | | | 107.2 86.36 | | 258.95 229.11 | 228.53 125.06 | 22.05 16.54 | | 388.34 151.37 44.56 22.01 | 53 16.12 | | |
|---------------------|---------------|-----------------|-----------------------|-----------|--------------|--------------|----------------|------------------|---------|---------------|--|---|--------------------------|---|-----------------------|--------------------------|---|
| normal no | set 2: se | Mean Me | | | 126.25 10 | | | 119.96 10 | | 349.4 25 | 219.26 22 | 28.48 22 | | 350.04 38 45.46 4 | 56.19 | | |
| metastati metastati | c: Std | Dev | | 8.38 | 79.49 | 6.58 | 34.83 | 10.95 | 10.8 | 79.07 | 30.07 | 7.51 | 31.83 | 23.09 33.75 | 14.93 | | |
| metastati | : <u>-</u> | Median | | 13.08 | 42.08 | 0.89 | 9.79 | 13.91 | 21.01 | 111.02 | 46.11 | 9.44 | 11.34 | 25.11 48.1 | 43.24 | | |
| | metastati | c: Mean | | 13.37 | 67.05 | 1.31 | 6.09 | 13.44 | 22.03 | 96.52 | 51.13 | 7.52 | 22.34 | 31.54 45.14 | 44.18 | | |
| | | p value | i . | 0 | #N/A | #N/A | 0 | 0.00142 | 0.00005 | 0.0263 | 0.00391 | #N/A | 0.00003 | 0.00005 #N/A | #N/A | | |
| fold | change | in metas | | 4.43 | #N/A | #N/A | 6.7 | 4.31 | 4.84 | 4.11 | 4.01 | #N/A | 21.37 | 9.56 #N/A | #N/A | | |
| | | Known Gene Name | Sec23 (S. cerevisiae) | homolog A | EST | TSE | EST | KIAA0096 protein | EST | EST | ficolin (collagen/fibrinogen domain-containing lectin) 2 (hucolin) | Zic family member 1 (odd- paired Drosophila homolog) | 1717 dihydropyrimidinase | sterol-C5-desaturase (fungal ERG3, delta-5-desaturase)- like EST | p53-responsive gene 2 | solute carrier family 23 | |
| | | Seq ID | | 1690 F | 1691 E | | | | 1706 | | 1709 (| 1716 | 1717 | 1750 H | 1758 | | |
| | | Genbank Seq ID | | D57823 | D57916 | D59294 | D59554 | D60769 | D61991 | D62103 | D63160 | D76435 | D78011 | D85181 D85433 | D86983 | | |
| | | Affy ID | | D57823_at | D57916 s at | rc D59294 at | rc D59554 f at | rc D60769 s at | | D62103_s_at | D63160_at | D76435 at | D78011_at | | _ D86983_at | | - |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| norma | set 2: | Std Dev | 66.7 | 415.69 | 589.37 | 168.21 | 96.66 | 79.24 | 31.82 | 27.37 | :88.21 | 165.09 | 138.17 | 83.35 26.24 31.96 |
|---------------------|-----------|-----------------|---|--|--------------|--------------|-------------------------|--------------|----------------|--------------|--------------|--------------|--------------------------------|--|
| normal n | | Median St | 176.16 | | 1040.46 5 | | 192.25 | 130.28 | 57.03 | 55.78 | 116.41 2 | 276.87 1 | | 93.71 6 71.82 5 52.54 6 |
| L | | | | | | | • | · | | | | | | |
| norma | set 2: | Mean | 167.38 | 712.98 | 1011.12 | 355.75 | 191.04 | 147.64 | 58.85 | 55.49 | 226.47 | 257.05 | 240.65 | 106.49 .64.94 .64.3 |
| metastati | c: Std | Dev | 32.5 | 8.02 | 62.74 | 34.39 | 28.07 | 33.97 | 28.12 | 23.17 | 20.72 | 90.09 | 12.33 | 13.56 8.75 22.72 |
| metastati metastati | ដ ់ | Median | 18.92 | 12.24 | 18.88 | 123.28 | 33.99 | -6.46 | 8.26 | 20.58 | 6.51 | 108.46 | 13.06 | 34.01 25.03 44.45 209.96 |
| | metastati | c: Mean | 5.13 | 8.24 | 42.65 | 138.06 | 40.92 | 13.91 | 12.72 | 9.35 | 6.98 | 111.86 | 19.64 | 36.92 22.16 44.91 233.85 |
| | | p value | 0 | 0.00002 | 0.00018 | #N/A | 0.00242 | 0.00088 | #N/A | #N/A | 0.02555 | #N/A | 0.0001 | #N/A #N/A #N/A 0.00025 |
| fold | change | in metas | 7.06 | 27.29 | 23.96 | #N/A | 4.01 | 4.79 | #N/A | #N/A | 4.36 | #N/A | 8.13 | #N/A #N/A #N/A 4.19 |
| | : | Known Gene Name | N-acetyltransferase 2 (arylamine N- 1767 acetyltransferase) | carbamoyl-phosphate synthetase 1, mitochondrial | EST. | EST | acyl-Coenzyme A oxídase | EST | EST | EST | EST | EST | dual specificity phosphatase 6 | butyrobetaine (gamma), 2- oxoglutarate dioxygenase (gamma-butyrobetaine hydroxylase) KIAA0541 protein EST |
| | ! | Sed ID |)) 1767 s | | 1792 E | 1794 E | | 1801 E | | | | 1810 E | 1814 | 1816 H 1817 H 1818 E |
| | | Genbank | D90042 | D90282 | F04611 | F04677 | F04944 | F09350 | F09687 | F09729 | F09979 | F10149 | F10276 | F10380 F10381 F10418 F10874 |
| | ! | Affy ID | D90042_at | D90282_at | rc_F04611_at | rc_F04677_at | rc_F04944_s_at | rc_F09350_at | rc_f09687_s_at | rc_F09729_at | rc_F09979_at | rc_F10149_at | rc_F10276_s_at | rc_F10380_at rc_F10381_s_at rc_F10418_at rc_F10874_f_at |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affy ID | Genbank | Seq ID | Known Gene Name | fold change in metas | p value | metastati c: Mean | metastati metastati c: c: Std Median Dev | metastati c: Std Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|---|--|--|--|---------------------------------------|---|---------------------------------|--|---|--|--|--|
| rc_F13624_at | F13624 | b 1825 ге | breast cancer anti-estrogen 1825 resistance 3 | #N/A | #N/A | 27.76 | 19.2 | 32.49 | 8.06 | 73.79 | 59.27 |
| rc_F13782_s_at | F13782 | 1827 L | 1827 LIM binding domain 2 | 4.17 | 0.00109 | -0.64 | 5.61 | 22.4 | 100.8 | 119.01 | 55.07 |
| rc_H01059_i_at rc_H02855_at | H01059 H02855 | 1830 tr 1832 E | solute carrier family 16 (monocarboxylic acid transporters), member 4 EST | #N/A 5.96 | #N/A 0.00458 | 9.13 5.06 | , 9.22 1.15 | 13.53 | 32.73 261.57 | 34.27 93.25 | 21.05 |
| rc_H03348_at rc_H05970_at | H03348 | 1833 c | laudin 1 ST | 5.77 #N/A | 0.0001 #N/A | 1.48 | 7.75 | 17.79 | 135.1 68.16 | 134.05 | 75.43 |
| rc_H09331_f_at rc_H09353_at | H09331 H09353 H09364 | 1865 E | EST EST EST | #N/A 23.06 | #N/A 0.00094 | 313.2 | 309.79 18.95 | 122.67 65.29 | 512.04 782.6 | 486.42 824.74 | 149.27 226.28 |
| 1 0000 1 00000 1 0000 1 0000 1 0000 1 0000 1 0000 1 0000 1 0000 1 0000 1 00000 1 0000 1 0000 1 0000 1 0000 1 0000 1 0000 1 0000 1 0000 1 00000 1 0000 1 0000 1 0000 1 0000 1 0000 1 0000 1 0000 1 0000 1 00000 1 0000 1 000 | | | | r r | 0.0020 | 5. | 2 | 2. | 25.75 | 9 | |
| rc_H11739_s_at | H11739 | 9 (6 d 1876 c | glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage system protein P) | 10.33 | 0.00023 | -21.2 | -37.69 | 28.32 | 290.05 | 210.37 | 244.67 |
| rc_H12593_at rc_H16098_at rc_H17472_s_at H19089_at rc_H19504_f_at | H12593 H16098 H17472 H19089 H19504 | 1885 EST 1888 EST 1894 EST 1895 EST | zinc-finger protein 265 EST EST EST | 10.72 #N/A #N/A #N/A 3.13 | 0.0056 #N/A #N/A #N/A 0.04948 | 39.96 45.29 12.2 76.83 | -9.15 37.73 9.54 44.82 49.48 | 111.74 26.92 8.57 89.14 64.05 | 515.58 87.28 31.08 76.54 227.6 | 506.07 94.49 31.66 74.4 221.91 | 224.5 38.15 13.28 40.21 125.23 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|----------------|----------------|----------|----------------------------|-------------|---------|-----------|---------------------|-----------|---------|---------|---------|
| | | | | change | | metastati | ິວ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank Seq ID | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| | | | | | | | | | | | |
| rc_H20543_at | H20543 | 1897 DI | 1897 DKFZP586B1621 protein | 31.03 | 0.00074 | 62.11 | 36.98 | 80.24 | 1684.92 | 1680.81 | 731.79 |
| H20627_at | H20627 | 1898 ES | ST | #N/A | #N/A | 30.59 | 30.04 | 23.84 | 79.8 | 87.46 | 43.37 |
| rc_H24081_at | H24081 | .1901 KI | AA1035 protein | #N/A | #N/A | 19.72 | 18.46 | 5.69 | 24.15 | 27.08 | 9.52 |
| rc_H25124_at | H25124 | 1903 ES | ST | 3.65 | 0.00004 | 20.47 | 14.4 | 12.59 | 95.09 | 85.36 | 34.91 |
| rc_H29568_at | H29568 | 1914 E | ST | 11.45 | 0.00058 | 232.54 | 140.09 | 218.85 | 2206.49 | 1884.76 | 783.84 |
| rc_H30270_at | H30270 | 1915 ES | EST | 17.09 | 0.00001 | 61.64 | 60.29 | 25.01 | 1224.59 | 1332.66 | 695.76 |
| H39627_at | H39627 | 1920 E | ST | #N/A | #N/A | 73.92 | 101.48 | 54.09 | 155.43 | 158.44 | 39.98 |
| | | 5 | tochrome P450, subfamily | | | | | | | | |
| H46990_at | H46990 | 1933 IIE | IIE (ethanol-inducible) | 3.2 | 0.00095 | -0.27 | -6.33 | 16.23 | 72.1 | 71.1 | 37.01 |
| | | ន | rboxypeptidase B2 | | | | | | | | |
| rc_H47838_at | H47838 | 1936 (p | (plasma) | 16.74 | 0.00002 | -26.99 | -41.82 | 25.87 | 401.55 | 412.78 | 188.81 |
| | | Ā | ATP-binding cassette, sub- | | | | | | | | |
| | | ā | family C (CFTR/MRP), | | | | | | | | |
| rc_H49417_s_at | H49417 | 1939 m | member 6 | #N/A | #N/A | 16.93 | 17.31 | 8.56 | 62.68 | 54.75 | 48.37 |
| rc_H52251_at | H52251 | 1942 ES | EST | #N/A | #N/A | 25.41 | 22.3 | 15.29 | 33.86 | 34.48 | 19.95 |
| rc_H53829_at | H53829 | 1946 ES | ST | #N/A | #N/A | 44.31 | 40.52 | 36.62 | 6.66 | 98.22 | 46.1 |
| | | | | | | | | | | | |
| | | 4 | 4-nitrophenylphosphatase | | ٠ | | | | | • | |
| 10 H56584 of | HEGERA | 4054 QC | domain and non-neuronal | u | c | 10.0 | 10 40 | 24 43 | 202 00 | 70 000 | 401 20 |
| rc_H56965_at | H56965 | 1952 ES | EST | 6.6 4/V# | A/N# | 0.18 | 0.68 | 12.9 | 61.03 | 73.89 | 30.12 |
| | | | • | | | 2 | 2 | ì | | 2 | - |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|--------------------------------|------------------|--------|-------------------------------------|--------------|--------------|-----------|---------------------|-----------|---------|---------|---------|
| | | • | | change | - | netactati | ë | S. Std | set 2. | set 2. | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| | | ū | protein phosphatase 2 | | | | | | | | |
| | | 0 | (formerly 2A), regulatory | | | | | | | | |
| rc H57850 at | H57850 | 1958 i | isoform | 3.02 | 0.00123 | 41.36 | 36.05 | 15.14 | 131.12 | 130.36 | 53.67 |
| 1 | | | formyltetrahydrofolate | | | | | | | | |
| rc h58692 s at | H58692 | 1960 c | dehydrogenase | 81.41 | 0 | -2.66 | 5.46 | 16.1 | 1886.69 | 1520.43 | 1129.35 |
| rc H60317 at | H60317 | 1965 E | EST | #N/A | #N/A | 18.86 | 22.34 | 11.81 | 38.92 | 39.1 | 18.67 |
| H61002 at | H61002 | 1967 E | EST SET | #N/A | #N/A | 96.89 | 23.38 | 102.25 | 88.87 | 93.79 | 36.34 |
| H66367_at | H66367 | | EST | 99.9 | 0.0001 | -5.44 | -2.13 | 18.66 | 155.38 | 149.31 | 70.77 |
| rc_H68239_f_at | H68239 | | EST | #N/A | #N/A | 26.96 | 17.94 | 15.81 | 92.99 | 66.18 | 51.67 |
| | | _ | v-Ki-ras2 Kirsten rat | | | | | | | | |
| | | 9) | sarcoma 2 viral oncogene | | | | | | | | |
| rc_H69138_at | H69138 | 1986 | homolog | 9.79 | 0.00142 | 42.88 | 30.87 | 38.35 | 328.51 | 313.13 | 218.8 |
| rc_H69565_at | H69565 | 1987 | : ST | 4.11 | 0.00002 | 15.45 | 16.92 | 9.4 | 92.6 | 94.91 | 42.92 |
| rc_H70485_at | H70485 | 1988 E | EST | #N/A | #N/A | 113.44 | 81.62 | 63.73 | 232.9 | 256.86 | 107.1 |
| rc H70554 at | H70554 | 1989 E | EST | 10.99 | 0 | -11.73 | -19.98 | 33.09 | 292.66 | 288.78 | 153.17 |
| rc_H70627_s_at | H70627 | 1990 E | EST | #N/A | #N/A | 17.89 | 17.62 | 8.85 | 42.72 | 41.34 | 11.7 |
| H72650_at | H72650 | 1994 E | EST SET | #N/A | #N/A | 81.21 | 98.79 | 41.38 | 122 | 125 | 42.24 |
| rc_H73535 s at | H73535 | 1996 E | EST | 68.9 | 0.00202 | 154.24 | 92.36 | 149.45 | 912.39 | 956.58 | 383.35 |
| rc H77494 at | H77494 | 1999 | EST | #N/A | #N/A | 9.65 | 6.64 | 15.53 | 42.67 | 47.73 | 27.54 |
| rc_H79820_at | H79820 | 2004 E | EST | 3.25 | 0.01466 | 10.64 | 7 | 22.28 | 113.3 | 113.19 | 92.96 |
| | | | | | | | | | | | |
| rc_H81070_f_at | H81070 | 2006 F | RNA helicase-related protein | 25.74 | 0.00126 | 93.43 | 62.01 | 121.29 | 2044.42 | 2051.11 | 615.84 |
| rc_H81964_s_at rc_H82424_at | H81964 H82424 | 2008 | SEC14 (S. cerevisiae)-like 1 EST | #N/A #N/A | #N/A #N/A | 9.95 | 15.7 | 12.54 | 29.9 | 21.14 | 20.56 |
| | | | | | |) | | : | | | :::: |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|----------------------------------|------------------|--------------|--|--------------------|--------------|----------------------|---------------------|----------------|------------------|------------------|-------------------|
| Affy ID | Genbank | Seq ID | Known Gene Name | cnange in metas | p value | metastati c: Mean | C: Median | C: Sta | set 2: Mean | set 2: Median | Set 2: Std Dev |
| rc_H82735_s_at | H82735 | 2010 | NOT3 (negative regulator of transcription 3, yeast) | #N/A | #N/A | -1.13 | 6.14 | 29.78 | 58.77 | 54.86 | 35.76 |
| rc_H82966_s_at rc_H83109_f_at | H82966 H83109 | 2011 2012 | apolipoprotein B (including Ag(x) antigen) EST | 3.42 16.55 | 0.00769 | 0.2 2.74 | 4.43 | 15.26 21.21 | 100.93 417.46 | 77.36 421.97 | 96.78 224.35 |
| rc_H87765_at | H87765 | 2017 | | 3.86 | 0.00131 | 2.5 | 7.91 | 19.6 | 94.26 | 93.78 | 60.64 |
| rc_H89980_at | H89980 | | | 31.13 | 0.00006 | 12.16 | 13.82 | . 12.3 | 990.48 | 1005.21 | 755.6 |
| rc_H90417_s_at rc_H93381_at | H90417 H93381 | 2028 2036 | EST | 4.17 24.23 | 0.015 0 | 14.23 6.08 | -10.79 -0.69 | 70.02 28.81 | 191.04 621.48 | 154.37 640.77 | 125.5 214.25 |
| , , | | | proline synthetase co- transcribed (bacterial | | | | | | | | |
| rc_H93562_at | H93562 | | homolog) | 3.17 | 0.00113 | 20.83 | 14.13 | 20.46 | 89.3 | 84.36 | 37.72 |
| rc_H93745_s_at | H93745 | 2040 | GS2 gene | #N/# | #N/A #// | 4.52 | 8.01 | 8.31 | 47.92 | 46.53 | 28.87 |
| rc_H95089_at | H95089 | | EST | Y/N# | YN# | 13.05 | 13.54 | 4.03 | 31.99 | 31.89 | 13.19 |
| rc_H95358_at | H95358 | 2049 | EST | 3.17 | 0.00182 | -5.85 | -5.95 | 11.57 | 74.74 | 72.66 | 42.36 |
| rc_H95978_at | H95978 | 2052 | EST | 8.55 | 0.00046 | 12.28 | 14.38 | 10.71 | 240.91 | 246.43 | 147.67 |
| rc_H96392_at | H96392 | 2053 | EST | #N/A | #N/A | 12.6 | 9.93 | 33.15 | 35.98 | 53.46 | 42.74 |
| rc_H96897_at | Н96897 | 2056 KIAA | KIAA0336 gene product | #N/A | #N/A | 35.22 | 19.98 | 42.05 | 48.46 | 50.39 | 33.36 |
| rc_H97587_s_at rc_H97670_at | H97587 H97670 | 2060 2061 | endothelin receptor type B EST | #N/A #N/A | #N/A #N/A | -5.07 23.31 | -3.35 29.71 | 14.02 31.35 | 39.65 101.35 | 36.54 128.08 | 21.7 56.78 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | fold | | | | | normal | normal | normal |
|------------------|---------|--|----------|----------------|------------|---|----------------|---------|---------|---------|
| | | | change | | metastatic | metastatic metastatic metastatic set 2: | metastatic | set 2: | set 2: | set 2: |
| Affy ID | Genbank | own Gene Name | in metas | p value : Mean | : Mean | : Median | : Std Dev Mean | Mean | Median | Std Dev |
| rc_H98083_at | H98083 | 2067 EST | 5.09 | 0.00025 11.73 | 11.73 | 11.11 | 3.97 | 121.51 | 104.88 | 70.52 |
| rc_H99393_s_at | H99393 | 2076 endothelin receptor type 3.43 | 3.43 | 0.00093 4.63 | 4.63 | 9.33 | 11.96 | 79.18 | 70.67 | 42.71 |
| | | | | | | | | | | |
| | | | | | | | | | | |
| | | | | | | | | | | |
| | | apolipoprotein A- | | | | | | | | |
| J00098_cds1_s_at | 100098 | 2086 I,apolipoprotein C-III corticosteroid binding | #N/¥ | #N/A | 96.71 | 29.74 | 269.46 | 9836.26 | 8356.89 | 8217.05 |
| J02943_at | J02943 | 2090 globulin | 18.98 | 0.00087 57.44 | 57.44 | 39.79 | 44.72 | 880.54 | 924.99 | 144.24 |
| | | insulin-like growth factor | | | | | | | | |
| J03242_s_at | J03242 | | 4.01 | 0.00042 40.37 | 40.37 | 45.32 | 15.53 | 179.86 | 172.23 | 104.64 |
| | | solute carrier family 2 (facilitated glucose | | · | | | | | | |
| J03810_at | J03810 | transporter), member 2 | 21.99 | 0.00004 21.12 | 21.12 | 16.81 | 24.34 | 571.66 | 593.24 | 118.08 |
| J04056_at | J04056 | | 5.19 | 0.00001 | 9.77 | 22.83 | 29.45 | 145.33 | 139.19 | 73.11 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|--------------------|---------|-------------|---|--------------------|---------|----------------------|---------------------|---------------|----------------|------------------|-------------------|
| . Affy ID | Genbank | Seq ID | Known Gene Name | change in metas | p value | metastati c: Mean | c: Median | c: Std Dev | set 2: Mean | set 2: Median | set 2: Std Dev |
| 10000 | . 070 | 24.08 | 1 DD alvacathrande | 18 02 | _ | 6 | 0 | 12.0 | 762 E1 | τ 7 |) 215.46 |
| J05037_at | J05037 | 2116 serine | serine dehydratase | 16.24 | 0.00015 | | 32.78 | 67.98 | 626.47 | 497.97 | 306.18 |
| J05158_at | J05158 | 2117 | carboxypeptidase N, 2117 polypeptide 2, 83kD | 8.52 | 0 | 2.27 | -7.78 | 21.43 | 209.63 | 218.19 | 81.01 |
| K02100_at | K02100 | 2123 | ornithine 2123 carbamoyltransferase | 10.24 | 0.00009 | 6.1 | 3.97 | 11.14 | 253.26 | 300.99 | 126.18 |
| K02402 | K02402 | 301C | coagulation factor IX (plasma thromboplastic component, Christmas | 28.81 | 0 00001 | 10.4 | 204 | 28.08 | 742 92 | 734 01 | 215 88 |
| K02766 at | KN9766 | 2126 com | complement component 9 | 21 24 | - | 23.15 | 14 02 | 21.18 | 652.66 | 656 14 | 320.1 |
| | |) ! | | . · ! |) | <u>}</u> | |) : : | | | |
| K03192 <u>f</u> at | K03192 | 2127 | cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 | 69.92 | 0 | 3.07 | 9.83 | 30.14 | 1767.25 | 1790.36 | 741.7 |
| K03192_f_at | K03192 | 2127 | cytochrome P450, subfamily IIA (phenobarbital-inducible), 7 polypeptide 6 | 69.92 | 0 | -1.32 | -6.49 | 12.52 | 1087.08 | 1034.79 | 451.92 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|-------------|----------------|--------|--|--------------------|---------|----------------------|---------------------|---------------|----------------|------------------|-------------------|
| Affy ID | Genbank Seq ID | Seq ID | Known Gene Name | change in metas | p value | metastati c: Mean | c: Median | c: Std Dev | set 2: Mean | set 2: Median | set 2: Std Dev |
| | | | cytochrome P450, subfamily I (aromatic compound- | | | | | | | | |
| L00389_f_at | L00389 | 2132 | inducible), polypeptide 2 | #N/A | #N/A | 69.38 | 86.39 | 64.81 | 222.59 | 184.81 | 121.16 |
| L00972_at | L00972 | 2133 | 2133 cystathionine-beta-synthase | 7.19 | 0.00008 | 30.44 | 29.08 | 21.44 | 268.9 | 253.33 | 153.24 |
| L04751_at | L04751 | 2138 | cytochrome P450, subfamily IVA, polypeptide 11 | 36.79 | 0.00004 | 32.39 | 23.29 | 32.64 | 1164.28 | 1183.4 | 334.08 |
| L07956 at | L07956 | 2148 | glucan (1,4-alpha-), branching enzyme 1 (glycogen branching enzyme, Andersen disease, glycogen storage disease type IV) | က် | 0.00029 | 0. 2. 2. 8. | 18.57 | 16.75 | | 184 57 | . 75 67 |
| 1 | | | 7 | | | 2 | 5 | 2 | - | 2 | 5 |
| L09717_at | L09717 | 2153 | lysosomal-associated membrane protein 2 | 4.06 | 0.00034 | 16.78 | 9.78 | 21.83 | 115.12 | 111.26 | 57.66 |
| L11244_s_at | L11244 | 2155 | complement component 4- binding protein, beta | 43.33 | 0 | 0.49 | -8.84 | 36.95 | 1147.16 | 1114.71 | 441.14 |
| L11244_s_at | L11244 | 2155 | complement component 4- binding protein, beta serine | 43.33 | 0 | -13.27 | -16.78 | 12.92 | 292.9 | 341.12 | 114.93 |
| L11931_at | L11931 | 2159 | hydroxymethyltransferase 1 (soluble) | 7.27 | 0.00041 | -18.81 | -31.65 | 26.43 | 188.25 | 181.11 | 114.89 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | netastati | normal | normal | normal |
|------------------------|------------------|----------------------------|--|--------------------|-----------------|----------------------|---------------------|----------------|-----------------|------------------|-------------------|
| Affy ID | Genbank Seg ID | Seq ID | Known Gene Name | change In metas | p value | metastati c: Mean | c: Median | c: Std Dev | set 2: Mean | set 2: Median | set 2: Std Dev |
| L13689_at | L13689 | 2164 | murine leukemia viral (bmi- 2164 1) oncogene homolog | #N/A | W/A | 32.33 | 34.76 | 18.57 | 63.42 | 77.84 | 36.39 |
| L19871_at | L19871 | 2172 | activating transcription factor 3 | #N/A | #N/A | 72.2 | 31. | 78.72 | 98.22 | 85.66 | 61.41 |
| L26336_at L27050_at | L26336 L27050 | 2185 heat s 2186 apolip | heat shock 70kD protein 2 apolipoprotein F | #N/A 10.26 | #N/A 0.00026 | 15.67 -23.8 | 14.08 -20.7 | 10.72 | 31.12 258.1 | 33.86 252.65 | 10.1 124.08 |
| L27841_at L32140_at | L27841 L32140 | 2187 2192 | pericentriolar material 1 afamin | #N/A 17.31 | #N/A 0.00003 | 58.75 9.44 | 58.95 16 | 40.61 10.75 | 90.06 426.89 | 82.71 464.97 | 28.43 221.11 |
| L32179_at | L32179 | 2193 | arylacetamide deacetylase (esterase) | 23.83 | . 0 | 22.94 | 24.96 | 16.06 | 688.97 | 675.71 | 305.75 |
| L32961_at | L32961 | 2194 | 4-aminobutyrate aminotransferase | #N/A | #N/A | 15.64 | 7.44 | 15.63 | 76.44 | 79.62 | 26.55 |
| L33799_at | L33799 | 2196 | procollagen C- endopeptidase enhancer | #N/A | #N/A | 87.45 | 73.5 | 89.14 | 230.54 | 248.1 | 130.08 |
| L34081_at | L34081 | 2199 | bile acid Coenzyme A: amino acid N- acyltransferase (glycine N- choloyltransferase) | 11.96 | 0.00008 | 6.09 | 4.41 | 14.12 | 314.27 | 330.42 | 174.43 |
| L41067_at | L41067 | 2213 | nuclear factor of activated T-cells, cytoplasmic 3 | 4.96 | 0.00473 | 57.54 | 60.41 | 36.65 | 261.77 | 228.95 | 119.65 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|----------------|---------|------------|--|----------|----------|-----------|---------------------|-----------|---------|--------|---------|
| | | | | change | | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| 10077 | 14007 | | nuclear factor of activated T- | 9 | 0007 | i c | 9 | 77 | 1001 | 2 | c c |
| L4106/_at | L4106/ | 6122 | cells, cytopiasinic s | 08.4 | 0.0047.5 | 30.23 | 13.00 | 32.44 | 13.07 | 00.77 | 20.20 |
| L44424_at | L44424 | | UMP-CMP kinase | #N/A | #N/A | 23.79 | 18.2 | 26.85 | 27.57 | 43.3 | 36.92 |
| L48516_at | L48516 | 2220 | paraoxonase 3 | 22.21 | 0.00004 | 36.05 | 28.85 | 37.25 | 896.92 | 896.69 | 337.05 |
| M10058_at | M10058 | 2230 | asialoglycoprotein receptor 1 metallothionein 1F | 23.96 | 0 | -47.64 | -42.75 | 75.52 | 591.6 | 603.68 | 188.25 |
| M10943 at | M10943 | 2234 | | 3.88 | 0 | 53.15 | 52.73 | 8.81 | 217.65 | 186.71 | 86.73 |
| M11437_cds1_at | M11437 | 2238 | kininogen | 18.38 | 0.00006 | 20.84 | 10.7 | 36.84 | 535.53 | 539.53 | 186.97 |
| | | | angiogenin, ribonuclease, | | | | | | | | |
| M11567_rna1_at | M11567 | 2239 | RNase A family, 5 lecithin-cholesterol | 32.25 | 0.0001 | 2.5 | -18.19 | 58.03 | 1019.17 | 968.25 | 273.9 |
| M12625_at | M12625 | 2245 | acyltransferase | 4.55 | 0.01584 | 110.15 | 128.06 | 73.94 | 447.22 | 425.61 | 138.52 |
| | | | protein C (inactivator of coagulation factors Va and | | | | | | | | |
| M12712_s_at | M12712 | 2246 | VIIIa) | 7.37 | 0.01866 | 107.12 | 177.74 | 113.63 | 646.94 | 628.19 | 326.15 |
| M12759_at | M12759 | 2247 | EST | #N/A | #N/A | 27.23 | 21.91 | 28.99 | 97.65 | 103.06 | 53.91 |
| | | | kallikrein B plasma, | | | | | | | | |
| M13143_at | M13143 | 2249 | (Fletcher factor) 1 | 10.39 | 0.00019 | 38.55 | 37.19 | 32.81 | 402.05 | 441.15 | 122.27 |
| | | | | | | | | | | | |
| | | | coagulation factor VII | | | | | | | | |
| M13232_s_at | M13232 | 2251 | conversion accelerator) | 5.9 | 0.00014 | 43.46 | 42.59 | 18.46 | 248.65 | 275.82 | 70.94 |
| M13699_at | M13699 | 2253 cèrul | ceruloplasmin (ferroxidase) | 15.85 | 0.00012 | 36.21 | 37.43 | 44.51 | 688.89 | 638.74 | 329.28 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|-------------|----------------|-----------------|--|----------|---------|-----------|---------------------|-----------|---------|---------|---------|
| | | | | change | | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank Seq ID | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| M14091_at | M14091 | 2257 | 2257 thyroxin-binding globulin | 10.66 | 0.00024 | 1.76 | 2.84 | 6.6 | 269,26 | 284.66 | 119.28 |
| 44.4777 | 1 | 000 | glutathione S-transferase A2,glutathione S-transferase | *** | *** | Š | ! | 1 | | | . 6 |
| W14///_s_at | M14/// | 2263 A3 pyr⊔ | A3 pyruvate kinase, liver and | Y/N# | ₹ ¥ | -24.81 | -27.52 | 27.54 | 2842.35 | 2790.19 | 1048.69 |
| M15465_s_at | M15465 | 2266 | RBC | 6.1 | 0.00069 | 3.06 | 3.32 | 11.38 | 158.56 | 132.57 | 102.11 |
| M15656_at | M15656 | 2268 | bisphosphate | 99.66 | 0 | -44.61 | -66.97 | 59.8 | 2911.78 | 2539.75 | 1883.04 |
| M16594_at | M16594 | 2272 | 2272 glutathione S-transferase A2 | 73.21 | 0 | -3.62 | -9.74 | 18.15 | 1602.11 | 1787.28 | 655.41 |
| M16967_at | M16967 | 2275 | coagulation factor V 2275 (proaccelerin, labile factor) | 5.56 | 0.00047 | 22.71 | 26.49 | 30.11 | 204.85 | 248.05 | 88.92 |
| M16973_at | M16973 | 2276 | complement component 8, beta polypeptide | 22.75 | 0.00001 | 20.03 | 23.43 | 31.9 | 686.99 | 714.82 | 203.16 |
| M16974_s_at | M16974 | 2277 | | 49.47 | 0.00046 | 28.24 | -6.62 | 84.02 | 1843.82 | 1920.94 | 524.62 |
| M17466_at | M17466 | 2279 | (Hageman factor) | 9.76 | 0.00285 | 94.21 | 63.47 | 93.18 | 706.11 | 724.27 | 261.46 |
| M20218_at | M20218 | 2288 | coagulation factor XI (plasma thromboplastin antecedent) | 6.4 | 0.00004 | 38.74 | 45.83 | 14.62 | 247.85 | 259.47 | 69.46 |
| M20786_at | M20786 | 2290 alpha | alpha-2-plasmin inhíbitor | 16.95 | 0.00709 | 58.96 | -14.12 | 171 | 1074.45 | 1132.06 | 620.06 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|--------------------------|------------------|--------------|--|----------------|---------|----------------|---------------------|----------------|----------------|------------------|-----------------|
| | | | | change | _ | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| M22324_at | M22324 | 2295 | alanyl (membrane) aminopeptidase (aminopeptidase N, aminopeptidase M, microsomal aminopeptidase, CD13, | #N/A | A/N# | 224.68 | 25.21 | 471.53 | 365.81 | 363.02 | 134.32 |
| M23234_s_at | M23234 | 2299 | ATP-binding cassette, subfamily B (MDR/TAP), member 4 | 10.05 | 0 | -4.37 | -11.49 | 29.05 | 255.03 | 242.93 | 112.08 |
| M24283_at | M24283 | 2303 | intercellular adhesion molecule 1 (CD54), human rhinovirus receptor acyl-Coenzyme A | 3.19 | 0.04985 | 77.88 | 34.05 | 69.45 | 207.07 | 183.74 | 121.4 |
| M26393_s_at M29194_at | M26393 M29194 | 2309 2315 | | 16.27 12.18 | 0.00007 | 16.81 18.97 | 12.39 11.91 | 48.16 30.61 | 677.3 352.8 | 775.54 332.28 | 358.47 87.49 |
| M30185_at | M30185 | 2321 | cholesteryl ester transfer protein, plasma | 3.83 | 0.0013 | -37.15 | -29.18 | 28.09 | 93.07 | 86.04 | 61.35 |
| M30257_s_at M30269_at | M30257 M30269 | 2322 2323 | vascular cell adhesion molecule 1 nidogen (enactin) | 3.11 | 0.00064 | 17.33 | 13.46 26.36 | 10.06 | 78.1 96.8 | 82.04 94.16 | 35.74 42.52 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affy ID | Genbank | Seq ID | Known Gene Name | fold change in metas | p value | metastati c: Mean | metastati c: Median | metastati metastati c: c: Std Median Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|----------------|---------|--------|---|----------------------------|---------|----------------------|---------------------------|--|--------------------------|----------------------------|-----------------------------|
| M33317_f_at | M33317 | 2338 | cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 7 | 32.63 | 0 | 44.53 | 35.06 | 29.77 | 1613.89 | 1437.34 | 1086.43 |
| M61853_at | M61853 | 2369 | cytochrome P450, subfamily IIC (mephenytoin 4- hydroxylase), polypeptide 18 | 7.82 | 0.00024 | 12.17 | 12.44 | 4.36 | 191.94 | 204.27 | 91.68 |
| M61854_s_at | M61854 | 2370 | cytochrome P450, subfamily IIC (mephenytoin 4- hydroxylase) | 3.3 | 0.04185 | 52.7 | 55.56 | 44.01 | 321.6 | 166.36 | 430.71 |
| M63967_at | M63967 | 2378 | 2378 aldehyde dehydrogenase 5 | 4.04 | 0.00058 | 34.59 | 36.64 | 5.46 | 164.38 | 178.74 | 87.39 |
| M64174_at | M64174 | 2379 | | #N/A | #N/A | 10.42 | 10.81 | 15.82 | 59.51 | 59.38 | 27.77 |
| M64554_rna1_at | M64554 | 2380 | coagulation factor Alli, b polypeptide | 5.87 | 0.00011 | -4.73 | -4.91 | 7.92 | 137.26 | 136.8 | 72.05 |
| M64590_at | M64590 | 2381 | glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine 2381 cleavage system protein P) | 6.41 | 0.00002 | 14.28 | 17.87 | 8.65 | 152.26 | 127.72 | 82.96 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affy ID | Genbank Seq ID | Seq ID | Known Gene Name | fold change in metas | p value | metastati c: Mean | metastati c: Median | metastati metastati c: c: Std Median Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|--------------------------|------------------|--------------|---|----------------------------|-----------------|----------------------|---------------------------|--|--------------------------|----------------------------|-----------------------------|
| M64929_at | M64929 | 2383 | protein phosphatase 2 (formerly 2A), regulatory subunit B (PR 52), alpha isoform | #N/A | #N/A | 64.63 | 38.82 | 95.31 | 106.38 | 88.13 | 67.95 |
| M65134_s_at | M65134 | 2385 | 2385 complement component 5 protein C inhibitor | 12.01 | 0.00012 | 2.31 | 3.66 | 7.43 | 303.27 | 337.95 | 151.33 |
| M68516_rna1_at | M68516 | 2387 | | 20.54 | 0 | -1.34 | -11.78 | 20.54 | 466.81 | 481.06 | 119.75 |
| M68840_at | M68840 | 2388 | 2388 monoamine oxidase A | 3.96 | 0.01396 | 42.09 | 28.9 | 45.52 | 182 | 201.64 | 94.09 |
| M69177_at M74719_at | M69177 M74719 | 2392 2396 | | 11.64 #N/A | 0.00001 #N/A | 21.97 8.97 | 20.41 15.15 | 20.83 | 322.15 46.89 | 340.27 | 94 12.66 |
| M75106_at | M75106 | 2397 | carboxypeptidase bz 2397 (plasma) | 40.63 | 0 | 19.77 | 30.19 | 22.24 | 1170.81 | 1124.59 | 329.77 |
| M76665_at M83216_s_at | M76665 M83216 | 2398 2407 | hydroxysteroid (11-beta) dehydrogenase 1 caldesmon 1 | 19.22 4.27 | 0.00004 | 33.41 7.89 | 17.51 2.98 | 23.86 | 628.97 105.21 | 670.88 109.38 | 196.22 55.62 |
| M83772_at | M83772 | 2409 | | 19.54 | 0 | 6.88 | -0.95 | 15.39 | 504.14 | 595.51 | 245.03 |
| M86873_s_at | M86873 | 2414 | plasminogen,plasminogen- like | 17.54 | 0 | -3.92 | -8.08 | 6.71 | 410.11 | 403.69 | 218.51 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati | metastati metastati | normal | normal | normal |
|--------------------------------|------------------|--------------|---|---------------|---------|--------------|----------------|---------------------|-------------------|-------------------|-------------------|
| ; | | | | change | _ | metastati | ö | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Sed ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| | | | | | | | | | | | |
| | | | SWI/SNF related, matrix associated, actin dependent | | | | | | | | |
| M88163_at | M88163 | 2416 | regulator of chromatin, subfamily a, member 1 | 3.37 | 0.00098 | 17.39 | 17.52 | 7.25 | 84.41 | 91.86 | 41.42 |
| M88338_at | M88338 | 2417 | serum constituent protein | #N/A | #N/A | 134.22 | 128.76 | 53.79 | 155.51 | 140.27 | 70.28 |
| M88468_at | M88468 | 2418 | mevalonate kiriase (mevalonic aciduria) methylmalonate- | #N/A | #N/A | 89.85 | 108.13 | 54.23 | 214.81 | 260.33 | 92.98 |
| M93405_at | M93405 | 2424 | semialdehyde dehydrogenase | 23.06 | 0 | 16.51 | 27.77 | 20.66 | 679.66 | 734.58 | 307.25 |
| M94065_at | M94065 | 2425 | alnydroorotate dehydrogenase | 6.47 | 0.00013 | 30.26 | 20.14 | 17.64 | 202.13 | 183.44 | 86.43 |
| M95767_at | M95767 | 2432 | 2432 chitobiase, di-N-acetyl- | 4.94 | 0.00004 | 26.01 | 37.13 | 21.53 | 173.07 | 153.03 | 92 |
| | | | CD36 antigen (collagen type | | | | | | | | |
| M98399 s at | M98399 | 2437 | I receptor, thrombospondin receptor) | #N/A | #N/A | 4.71 | 4 58 | rc C | 47.65 | 38 28 | 35.2 |
| rc_N21626_at | N21626 | | EST | W/A | #N/A | 29.77 | 24.63 | 25.09 | 59.18 | 46.15 | 33.56 |
| rc_N22404_at | N22404 | 2450 | EST | 3.99 | 0.01152 | 30.59 | 23.97 | 55.32 | 193.63 | 130.09 | 162.83 |
| rc_N22434_at | N22434 | 2451 | EST | 4.37 | 0.01725 | 100.96 | 66.55 | 110.62 | 334.51 | 328.98 | 150.7 |
| rc_N22938_s_at rc_N24879_at | N22938 N24879 | 2453 2460 | serum amyloid A4, constitutive EST | 35.39 9.44 | 0.00128 | 163.4 6.2 | 145.31 5.49 | 160.24 23.52 | 4244.44 280.93 | 3330.35 337.77 | 2358.25 165.77 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | fold | | | metastati | metastati metastati | normal | normal | normal |
|--------------|---------|------------------------------|----------|-----------|-----------|-----------|---------------------|--------|----------------|---------|
| | | | change | 5 | metastati | : | Std. | Set 2: | set 2: | set 2: |
| Affy ID | Genbank | Sed ID Known Gene Name | in metas | p value (| c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc_N24973_at | N24973 | 2462 EST | #N/A | #N/A | 30.46 | 28.45 | 11.29 | 56.57 | 59.09 | 22.87 |
| rc N25262 at | N25262 | 2465 EST | #N/A | #N/A | 31.16 | 28.04 | 9.03 | 64.13 | 64.97 | 18.29 |
| rc_N27524_at | N27524 | 2472 EST | 3.13 | 0.00216 | 14.18 | 22.08 | 20.51 | 87.67 | 98.26 | 42.62 |
| N27670 at | N27670 | progesterone membrane | 6 15 | 0.00321 | 38 66 | 15.48 | 47 19 | 252 94 | 292.34 | 133.48 |
| ואביסיטים | 0/0/70 | | 3 | 0.0032 | 20.00 |) - | | £0.707 | 4 36.01 | 7.00 |
| | | kynurenine 3- | | | ٠ | | | | ٠ | |
| | 1 | mon | į | | | 1 | • | i | | 4 |
| rc_N29353_at | N29353 | | 4.78 | 0.00019 | -4.7 | -5.88 | 13.94 | 109.79 | 101.27 | 53.38 |
| rc_N29543_at | N29543 | EST | #N/A | #N/A | 11.66 | 9.07 | 5.04 | 59.21 | 51.22 | 30.43 |
| rc_N29740_at | N29740 | | #N/A | #N/A | 48.71 | 27.45 | 46.9 | 80.01 | 6.77 | 37.18 |
| rc_N31952_at | N31952 | 2490 EST | 3.13 | 0.01481 | 36.21 | 27.5 | 25.04 | 133.49 | 147.66 | 83.03 |
| rc_N32071_at | N32071 | | 8.75 | 900000 | 40.89 | 43.12 | 15.37 | 421.21 | 488.49 | 238.4 |
| | | | | | | | | , | | |
| rc_N34804_at | N34804 | | 8.08 | 0.00028 | 6.99 | 5.94 | 33.73 | 344.08 | 176.4 | 532.11 |
| rc_N34919_at | N34919 | | #N/A | #N/A | 7.39 | œ | 6.82 | 52.96 | 44.23 | 24.23 |
| rc_N35247_at | N35247 | | #N/A | #N/A | 39.9 | 33.56 | 15.03 | 117.08 | 116.14 | 57.05 |
| rc_N36085_at | N36085 | | #N/A | #N/A | 20.98 | 19.21 | 13.89 | 58.28 | 28 | 22.95 |
| N40320_at | N40320 | 2514 EST | 7.56 | 0.01584 | 99.72 | 83.43 | 128.45 | 558.37 | 595.16 | 250.67 |
| rc_N45320_at | N45320 | | #N/A | #N/A | 11.39 | 11.26 | 2.06 | 50.75 | 36 | 39.19 |
| rc_N45998_at | N45998 | 2520 EST | 3.14 | 0.00337 | 28.42 | 22.62 | 13.52 | 99.42 | 110.53 | 49.55 |
| rc_N47469_at | N47469 | | 3.34 | 0.00329 | 20.35 | 13.91 | 20.65 | 104.74 | 99.87 | 61.84 |
| rc_N48155_at | N48155 | 2525 EST | #N/A | #N/A | 17.08 | 16.77 | 14.66 | 97.37 | 54.69 | 2.66 |
| | | odentor-related protein | | | | | | | | |
| rc N48315 at | N48315 | 2527 complex 2, mu 1 subunit | 5.3 | 0.0149 | 62.03 | 30.66 | 52.29 | 368.37 | 331.28 | 260.93 |
| rc_N48674_at | N48674 | 2530 EST | 4.06 | 0.00028 | 2.7 | 4.57 | 10.52 | 94.61 | 72.82 | 54.62 |
| rc_N50038_at | N50038 | | #N/A | #N/A | 0.25 | 5.19 | 45.95 | 47.04 | 41.45 | 16.56 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | fold | | | metastati | metastati metastati | normal | normal | normal |
|--------------------------------|------------------|---|---------------|--------------------|----------------|----------------|---------------------|-------------------|------------------|-------------------|
| | | | change | _ | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID Known Gene Name | In metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc N51117 at | N51117 | 2544 EST | 9.68 | 0.00081 | 8.23 | 4.5 | 11.88 | 327.53 | 296.4 | 266.42 |
| rc N52845 at | N52845 | 2554 EST | 5.53 | 0.00088 | 28.56 | 30.14 | 14.95 | 219.38 | 230.31 | 149.67 |
| rc_N53757_at | N53757 | 2560 EST | 3.97 | 0.00255 | 99.49 | 80.7 | 56.09 | 365.45 | 324.21 | 118.42 |
| | | | | | | | | | | - |
| rc_N54053_at rc_N54311_at | N54053 N54311 | 2561 24kD 2564 EST | 60.39 | 0.00087 | 44.3 9.77 | -18.16 9.36 | 166.47 25.17 | 2749.73 160.29 | 2790.77 | 1230.33 124.7 |
| rc_N54604_at | N54604 | 2570 EST | 3.72 | 0.00741 | 2.36 | 2.08 | 35.23 | 133.06 | 119.1 | 109.95 |
| | | cytidine monophosphate-N- acetylneuraminic acid hydroxylase (CMP-N- | | | | | | | | · |
| rc_N54792_at rc_N55272_at | N54792 N55272 | acetylneuraminate 2571 monooxygenase) 2574 EST | 3.23 #N/A | 0.00099 #N/A | 28.36 | 27.42 13.6 | 6.39 24.4 | 110.19 45.08 | 77.69 | 87.32 14.77 |
| | | CCAAT/enhancer binding | | | 6 | ; | 9 | 9 | . (| |
| N57464_at | N57464 | 2576 protein (C/EBP), deita formiminotransferase | 14.69 | 0.00018 | 22.28 | 11.43 | 38.78 | 429.29 | 442.58 | 169.83 |
| rc_N57934_s_at rc_N59089_at | N57934 N59089 | 2577 cyclodeaminase 2581 EST | 13.81 | 0.00171 0.00055 | 806.55 38.4 | 477.9 36.58 | 939.39 24.6 | 9051.2 215.54 | 6373.16 167.6 | 8542.52 133.64 |
| | | aminomethyltransferase (glycine cleavage system | | | | | | | | |
| rc_N59532_s_at | N59532 | 2585 protein T) | 6.73 | 0.00005 | 12.43 | 2.55 | 17.66 | 196.37 | 155.84 | 122.13 |
| rc_N59543_at rc_N59550_at | N59543 N59550 | 2587 PDZ domain containing 1 2588 EST | 9.96 25.56 | 0.00052 0.00024 | 1.2 19.9 | -2.89 2.53 | 17.29 65.91 | 303.49 855.88 | 259.23 774.72 | 225.96 358.3 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | 1 | | fold | | metastati | metastati metastati | metastati c: Std | normal set 2: | normal set 2: | normal set 2: |
|----------------|----------------|-----------|------------------------------|----------|---------------|---------------|---------------------|---------------------|------------------|------------------|------------------|
| Affy ID | Genbank Seq ID | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc N62523 at | N62523 | 2592 h | hepatic leukemia factor | 5.02 | 0.00087 | 9.04 | 6.67 | 16.46 | 144.8 | 148.62 | 85.03 |
| rc_N62922_at | N62922 | 2596 E | EST | #N/A | #N/A | 14.92 | 14.15 | 2.75 | 54.94 | 52.4 | 18.93 |
| rc_N63047_at | N63047 | 2597 E | EST . | #N/A | #N/A | 4.13 | 0.93 | 9.72 | 14.78 | 15.63 | 8.24 |
| | | J | cell division cycle 42 (GTP- | | | | | | | | |
| rc_N63172_at | N63172 | | binding protein, 25kD) | #N/A | #N/A | 26.88 | 33.1 | 24.75 | 60.99 | 57.64 | 40.01 |
| rc_N63698_at | N63698 | 2604 | EST | 7.92 | 0.00001 | 19.14 | 18.14 | 12.14 | 221.25 | 212.4 | 122.32 |
| rc N64017 at | N64017 | | EST | 5.27 | 0.00022 | 5.45 | 8.71 | 8.71 | 122.34 | 135.27 | 55.58 |
| rc_N64436_at | N64436 | | EST | 3.51 | 0.00441 | 24.46 | 20.38 | 21.08 | 119.24 | 133.44 | 99.02 |
| rc_N65959_at | N65959 | | EST | 5.43 | 0.00044 | 3.51 | 3.25 | 22.24 | 142.87 | 149.26 | 79.26 |
| | | | | | | | | | | | |
| | | | progesterone membrane | o o | | c c | Č | 0 | 740 | 707 | 7077 |
| rc_N66130_at | N66130 | 2614 [| oinding protein | 3.58 | 0.0.0 4.14 | 29.02 | 27.05 | 33.88 | 62 58 | 50 13 | 51 00 |
| rc_N66613_at | N66613 | 71.97 | - 011 | ¥2. | ₹ ¥ | 10.02 | 46.27 | 20.33 | 02:30 | 2.00 | 6 |
| | | | | | | | | | | | |
| | | _ | prion protein (p27-30) | | | | | | | | |
| | | - \ | (Creutzfeld-Jakob disease, | | | | | | | | |
| | | - • | Gerstmann-Strausier- | | | | | | | | |
| | | | | i i | 0 | | | 7 | 000 | 00,404 | 1 |
| rc_N67009_s_at | N67009 | | familial insomnia) | 3.55 | 0.00956 | - | 11.35 52.5 | 38.17 | 22.221 | 124.39 | 60.00 |
| rc_N67105_at | N67105 | 2624 | ESI | 5.01 | 0.00176 | <i>و۲.</i> رہ | U.Z.1 | 10.38 | 143.30 | 102.9 | 4.1.4 |
| | | | inentin-like arowth factor 1 | | | | | | | | |
| rc N67876 c of | N67876 | mos) 8080 | (somatomedin C) | 8 89 | 0.00042 | 1181 | 15.88 | 8 82 | 267.41 | 145.79 | 238.35 |
| ור ויסיסים | 0.00.001 | 7070 | | 8 | 2000 | - - | | | | | |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | · | metastati | metastati c: | metastati metastati c: c: Std | normal set 2: | normal set 2: | normal set 2: |
|-----------------------------|------------------|--------------|--|--------------|-------------|---------------|-----------------|----------------------------------|------------------|------------------|------------------|
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| Te N67974 | A5079N | 0830 | activating transcription factor 4 (tax-responsive enhancer | ∇ /N# | ∇/N# | 787 | - C | 15.07 | . 68 | 74 13 | 50.47 |
| rc_N68133_at | N68133 | 2633 | EST EST | W/W# | #N/A | 2.35 | -5.53 | 16.59 | 58.69 | 37.21 | 57.82 |
| rc N68596 s at | Neason | 2636 | betaine-homocysteine methyltransferase | 40 99 | c | 77 | 13 84 | 10 15 | 1061.81 | 1013 27 | 574.42 |
| rc_N68871_at | N68871 | 2638 | EST | #N/A | #N/A | 4.3 | 4.02 | 8.66 | 34.08 | 36.71 | 14.33 |
| rc_N70305_at | N70305 | 2655 | EST | 4.41 | 0.00078 | -13.99 | -13.48 | 10.76 | 105.8 | 97.72 | 72.15 |
| rc_N70358_s_at | N70358 | 2657 | growth hormone receptor kidney- and liver-specific | 34.35 | 0 | 20.93 | 29.26 | 36.06 | 1449.57 | 1157.97 | 1115.22 |
| rc_N71542_at | N71542 | 2665 | gene | 21.05 | 0 | -14.76 | -18.28 | 14.3 | 471.65 | 486.43 | 226.63 |
| rc_n71935_s_at | N71935 | 2667 | multiple PDZ domain protein | 16.28 | 0 | 12.84 | 19.17 | 14.1 | 410.92 | 422.1 | 218.42 |
| rc_N72200_at | N72200 | 2669 | EST | #N/A | #N/A | 12.93 | 14.15 | 14.26 | 57.8 | 56.14 | 30.12 |
| rc_N73561_at | N73561 | 2676 | EST | 12.96 | 0.00011 | 17.04 | 4.48 | 25.06 | 516.79 | 458.57 | 393.69 |
| rc_N73883_at | N73883 | 2682 | EST | 9.35 | 0 | 18.47 | 23.48 | 17.67 | 256.78 | 276.41 | 95.34 |
| rc N74025 at | N74025 | 2685 | deiodinase, iodothyronine, type I | 22.79 | 0 | -7.64 | 2.14 | 15.95 | 527.67 | 515.68 | 256.99 |
| rc_N74422_at | N74422 | 2686 | EST | 30.32 | 0 | 23.57 | 28.41 | 36.93 | 1063.07 | 1094.48 | 371.26 |
| | | | collectin sub-family member | | | | | | | | |
| rc_N74624_at | N74624 | 2688 | 10 (C-type lectin) | #N/A | #N/A | 33.8 | 34.24 | 5.13 | 71.81 | 62.09 | 43.57 |
| rc_N75072_at | N75072 | 2689 | EST | 3.73 | 0.00379 | 5.14 | 6.21 | 15.5 | 104.12 | 104.99 | 81.79 |
| N75120_at rc_N76086_r_at | N75120 N76086 | 2690 2695 | EST . | AN# | 4/N# #/A | 20.59 8.23 | 23.42 7.19 | 10.65 6.54 | 61.42 55.55 | 58.88 55.02 | 13.84 35.47 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|----------------------------------|------------------|---------------|---|----------|-------------------|---------------|---------------------|---------------|-----------------|------------------|-----------------|
| | | | | change | | metastati | ၓ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc_N80129_i_at | N80129 | 2703 r | metallothionein 1L | 66.48 | 0.00415 | 179.8 | 104.5 | 643.06 | 10421.47 | 6358.88 | 9618.38 |
| N87590_at | N87590 | 2707 E | EST | #N/A | #N/A | 1.04 | -3.72 | 66.6 | 64.67 | 59.2 | 35.53 |
| rc N91882 at | N91882 | | alpha2,3-sialyltransferase | 14.52 | 0.00024 | -0.18 | 0.18 | 34.36 | 512.3 | 550.51 | 320.62 |
| rc_N93191_at | N93191 | 2730 E | EST | 3.25 | 0.00232 | 174.54 | 168.53 | 56.36 | 659.95 | 565.08 | 409.15 |
| rc_N93403_at | N93403 | | EST | #N/A | #N/A | 4.44 | 7.63 | 11.8 | 48.84 | 38.39 | 30.93 |
| rc_N93470_at | N93470 | | EST | #N/A | #N/A | 14.01 | 8.69 | 17.09 | 77.86 | 66.84 | 70.45 |
| rc_N93764_at | N93764 | | EST | 5.92 | 0.0109 | 113.46 | 68.13 | 96.98 | 771.74 | 792.59 | 504.88 |
| N94146 at | N94146 | | EST | 8.05 | 0 | 28.92 | 34.7 | 14.14 | 258.82 | 265.55 | 101.53 |
| N99542_at | N99542 | | orosomucoid 1 | 8.41 | 0.00001 | 6.74 | 10.14 | 21.89 | 226.06 | 187.68 | 146.11 |
| rc_R02365_f_at rc_R05309_f_at | R02365 R05309 | 2755 c 2759 E | glucose-6-phosphatase, catalytic (glycogen storage disease type I, von Gierke disease) | 3.72 | 0.00124 0.0008 | 0.92 21.94 | -5.3 13.92 | 9.39 18.75 | 768.72 96.81 | 444.03 100.54 | 841.02 31.73 |
| | | | SEC24 (S. cerevisiae) related gene family, member | | | | | | | | |
| rc_R05490_at | R05490 | | m | 3.03 | 0.00317 | 2.87 | 3.49 | 12.23 | 75.28 | 78.17 | 48.59 |
| rc_R06002_s_at | R06002 | | EST | 10.15 | 0.00003 | 10.52 | 8.26 | 20.8 | 292.89 | 264.17 | 160.28 |
| rc_R06271_s_at | R06271 | | EST | 6.14 | 0.00063 | 23.1 | 18.95 | 21.07 | 208.75 | 221.65 | 107.72 |
| rc_R06273_f_at | R06273 | | EST | W/V# | #N/A | 138 | 103.77 | 75.66 | 235.98 | 183.78 | 122.56 |
| rc_R06543_at | R06543 | 2769 E | EST | #N/A | #N/A | 61.75 | 61.42 | 88.77 | 163.57 | 144.04 | 96.98 |
| rc_R06746_at | R06746 | | EST | 15.77 | 0.0000 | 13.37 | -0.9 | 39.24 | 455.48 | 410.16 | 191.35 |
| rc_R06860_at | R06860 | | EST | #N/A | #N/A | 84.61 | 75.89 | 71.35 | 104.51 | 116.59 | 44.71 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|----------------|---------|----------------|-----------------------------|----------|---------|-----------|---------------------|-----------|---------|---------|---------|
| | | | | change | | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| | | <u>15</u> | glucokinase (hexokinase 4) | | | | | | | | |
| rc_R06977_f_at | R06977 | 2775 re | regulatory protein | 6.1 | 0.00049 | -47.7 | -38.97 | 37.79 | 152.22 | 150.69 | 87.15 |
| rc_R08564_at | R08564 | 2780 pla | asminogen-like | 60.18 | 0.00091 | 69.78 | 24 | 105.91 | 2485.63 | 2384.62 | 531.43 |
| rc_R09053_at | R09053 | | EST | 5.9 | 0.0002 | 6.45 | 17.17 | 35.01 | 186.87 | 204.61 | 113.19 |
| rc_R10287_at | R10287 | | ST | 11.2 | 0.0003 | 7.46 | 4.94 | 35.03 | 376.82 | 434.91 | 198.74 |
| rc R10684 at | R10684 | | ST | 5.51 | 0.00741 | 28.67 | 1.31 | 62.39 | 217.05 | 221.32 | 122.94 |
| rc_R12472_at | R12472 | | ST | 55.18 | 0.00011 | 32.21 | 3.13 | 76.26 | 2002.79 | 1930.03 | 1097.37 |
| rc_R12579_at | R12579 | | EST | 3.5 | 0.00137 | 54.49 | 37.84 | 26.65 | 198.7 | 194.59 | 92.03 |
| R19808_at | R19808 | | ST | #N/A | W/A | 9.95 | 12.2 | 25.55 | 27.53 | 25.43 | 10.61 |
| rc_R31104_f_at | R31104 | | . LS | #N/A | #N/A | -6.45 | 0.74 | 37.65 | 32.27 | 26.77 | 13.57 |
| R31641_at | R31641 | 2814 E | EST | 9.96 | 0.00011 | 29.02 | 21.83 | 31.18 | 350.89 | 330.68 | 166.29 |
| rc_R33146_at | R33146 | | ST | 7 | 0.00043 | 3.32 | 2.1 | 10.76 | 179.67 | 195.02 | 107 |
| | | | , | | | | | | | | |
| rc_R34362_at | R34362 | | KIAA0327 gene product | 5.4 | 0.04615 | 621.7 | 353.73 | 99.689 | 2107.55 | 2121.04 | 1000.75 |
| rc_R39238_at | R39238 | 2836 sy | synaptojanin 1 | #N/A | #N/A | 16.06 | 16.03 | 9.94 | 26.8 | 23.84 | 16.46 |
| | | | iecimin-cholesteroi | | | | | | | | |
| rc_R40395_s_at | R40395 | | acyltransferase | 37.33 | 0.00032 | 0.98 | -46.23 | 77.82 | 1192.03 | 1230.66 | 281.28 |
| rc_R40556_s_at | R40556 | 2843 E | ST | 3.57 | 0.00184 | 8.98 | 8.28 | 11.75 | 91.55 | 95.59 | 58.66 |
| | | ָ _ס | crystallin, zeta (quinone | | | | | | | | |
| rc_R40946_f_at | R40946 | | reductase) | 6.14 | 0.00156 | 18.65 | 19.52 | 14.3 | 205.48 | 220.3 | 137.27 |
| rc_R43174_s_at | R43174 | | paraoxonase 1 | 74.04 | 0.00038 | 47.94 | -8.83 | 119.68 | 2926.85 | 3104.07 | 1318.41 |
| | | - | downregulated in renal cell | | | | | | | | |
| rc_R43347_at | R43347 | 2849 ce | carcinoma | #N/A | #N/A | 36.93 | 32.68 | 30.41 | 105.66 | 100.62 | 28.62 |
| rc_R43365_at | R43365 | | EST | 5.37 | 0.00103 | 5.57 | 6.67 | 12.18 | 150.58 | 112.8 | 122.96 |
| rc_R43910_at | R43910 | 2852 E | ST | 5.71 | 0 | 23.41 | 24.6 | 11.1 | 156.7 | 166.34 | 52.21 |
| rc_R45480_i_at | R45480 | 2863 cy | cyclin K | #N/A | #N/A | 27.21 | 17.36 | 21.39 | 71.04 | 79.77 | 50.88 |
| rc_R49602_at | R49602 | 2885 E | EST | 36.5 | 0.00001 | 19.25 | 9.65 | 26.41 | 970.39 | 905.54 | 322.69 |
| rc_R51831_at | R51831 | | ST | 5.03 | 0.00761 | 5.89 | 2.5 | 44.67 | 221.54 | 195.97 | 179.99 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|----------------|---------|--------------------------|---|----------|-------------|-----------|---------------------|-----------|--------|--------|-------------|
| | | | | change | _ | metastati | ຮ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID Kn | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc R52822 i at | R52822 | 2896 EST | | 12.79 | 0.00001 | 10.78 | 0.35 | 28.51 | 380.07 | 441.17 | 180.12 |
| rc R53044 at | R53044 | 2898 KIAA09 | KIAA0981 protein | #N/A | #N/A | 17.84 | 25.74 | 11.45 | 48.77 | 44.34 | 18.43 |
| rc_R53891_at | R53891 | 2900 EST | | #N/A | #N/A | 9.88 | 5.59 | 10.68 | 66.28 | 52.16 | 49.21 |
| rc_R59312_at | R59312 | | | #N/A | #N/A | 34.37 | 35.83 | 22.13 | 94.44 | 91.55 | 36.35 |
| rc R59722 at | R59722 | | | 19.74 | 0.00016 | 18.78 | -4.94 | 51.34 | 628.69 | 642.78 | 316.88 |
| rc_R60959_at | R60959 | | | #N/A | #N/A | 31.21 | 23.86 | 17.79 | 72.94 | 69.87 | 45.25 |
| rc R62519 f at | R62519 | 2926 EST | | 10.83 | 0.00243 | 73.01 | 104.54 | 56.5 | 707.38 | 751.42 | 357.22 |
| R63545_at | R63545 | 2927 EST | | 5.4 | 0.00202 | 2.58 | 9.54 | 17.4 | 157.98 | 128.33 | 139.55 |
| to 167620 cz | De2724 | fatty-ac | fatty-acid-Coenzyme A | V/V# | V/W# | 200 | 47.43 | 10 17 | 72 67 | 30 C | 2 7 7 |
| R64534 at | R64534 | | EST | W/W# | V W # | 12.11 | 14.54 | 5.77 | 40.32 | 40.98 | 17.27 |
| l | | | | | | | | | | | |
| | , | kynure | enine 3- | | | | | | | | |
| rc R65593 s at | R65593 | 11101100 2935 (kynure | IIIoliouxygeriase (kynurenine 3-hydroxylase) | 21.27 | 0.00007 | 28.84 | 18.5 | 43.46 | 704.58 | 812.38 | 352.86 |
| 1 | | gamma | gamma-glutamyl | | | | | | | | |
| rc R67970 s at | R67970 | 2940 carboxylase | ylase | 9.45 | 0.00212 | 85.44 | 59.33 | 76.67 | 655.11 | 692.66 | 267.95 |
| rc R70319 at | R70319 | | | #N/A | #N/A | 5.34 | 2.91 | 8.92 | 29.23 | 29.18 | 10.78 |
| rc_R70791_at | R70791 | 2949 EST | | #N/A | #N/A | 127.11 | 110.15 | 41.83 | 73.19 | 57.2 | 56.17 |
| | | | | | | | | | | | |
| | | | differentially expressed in | ; | : | | | ! | ; | . ! | |
| R71459_at | R71459 | | adenocarcinoma of the lung | #N/# | W/A | 16.9 | 11.46 | 16.47 | 37.42 | 30.98 | 20.83 |
| rc_R73468_at | R73468 | 2957 EST | | #N/A | #N/A | 9.97 | 10.43 | 69.6 | 38.29 | 44.82 | 17.14 |
| rc_R73816_at | R73816 | | | 11.53 | 0.00259 | -26.88 | -30.68 | 10.09 | 462.31 | 374.67 | 450.91 |
| R77539_at | R77539 | 2965 EST | | #N/A | #N/A | 5.91 | 14.55 | 25.48 | 70.5 | 59.8 | 31.72 |
| rc_R78713_s_at | R78713 | 2968 EST | | #N/A | #N/A | 0.27 | -1.59 | 8.92 | 70.57 | 57.33 | 56.37 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | norma | normal |
|----------------|---------|---------|--|----------|---------|-----------|---------------------|-----------|---------|---------|---------|
| | | | | change | _ | metastati | ၓ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| | | đ | phosphatidylserine | | | | | | | | |
| R82229_at | R82229 | 2975 de | ecarboxylase | 3.08 | 0.03455 | 84.63 | 80.61 | 67.01 | 228.19 | 232.06 | 58.38 |
| rc_R91060_at | R91060 | 2981 E | ST | #N/A | #N/A | 59.14 | 37.15 | 56.04 | 85.88 | 61.99 | 68.89 |
| rc R93507 at | R93507 | 2991 E | ST | #N/A | #N/A | 19.53 | 20.08 | 7 | 67.62 | 67.56 | 44.22 |
| rc R93714 at | R93714 | 2992 fe | stuin B | 10.42 | 0.00043 | -60.74 | -65.14 | 57.99 | 280.62 | 301.36 | 203.86 |
| rc R93908 at | R93908 | 2994 E | EST | 5.8 | 0.02699 | 66.75 | 29.54 | 129.46 | 348.58 | 301.29 | 173.81 |
| rc R94674 s at | R94674 | 2996 E | ST | 10.76 | 0.00008 | 0.91 | 4.15 | 14.66 | 269.49 | 288.56 | 148.06 |
| rc_R96822_at | R96822 | 3000 E | ST . | 8.25 | 0.00008 | 16.45 | 17.01 | 30.67 | 250.81 | 257.18 | 131.85 |
| | | <u></u> | cytochrome P450, subfamily | | | | | | | | |
| rc R97419 at | R97419 | - | IIIB (sterol 12-alpha- ydroxylase), polypeptide 1 | 65.07 | 0.0039 | 152.16 | -3.42 | 302.56 | 4114.39 | 3837.4 | 1458.43 |
| rc_R97804_at | R97804 | 3008 E | EST | #N/A | #N/A | 144.46 | 187.67 | 92.8 | 193.05 | 201.89 | 61.84 |
| R98073_at | R98073 | 3009 E | ST | 46.87 | 0 | 4.33 | -9.52 | 11.43 | 1161.51 | 1291.95 | 564.26 |
| rc_R98105_at | R98105 | 3011 E | ST | #N/A | #N/A | 132.16 | 97.16 | 141.15 | 373.83 | 342.91 | 215.89 |
| rc_R98413_at | R98413 | 3012 E | ST | 9.71 | 0.00007 | 38.74 | 28.05 | 21.96 | 402.49 | 443.47 | 212.95 |
| rc_R98624_at | R98624 | 3013 E | ST | 21.32 | 0 | 15.01 | 4.48 | 22.97 | 686.81 | 542.22 | 466.97 |
| rc_R99014_at | R99014 | 3015 E | ST | #N/A | #N/A | 26.99 | 17.82 | 29.49 | 49.82 | 52.35 | 18 |
| | | O | CD5 antigen-like (scavenger | | | | | | | | |
| rc_R99591_at | R99591 | 3016 re | 3016 receptor cysteine rich family) | 9.52 | 0.00006 | 87.08 | 72.59 | 28.25 | 1035.39 | 904.92 | 691.83 |
| S62539_s_at | S62539 | 3022 in | 3022 insulin receptor substrate 1 | 3.7 | 0.01307 | 28.26 | 13.05 | 40.92 | 132.59 | 97.52 | 103.35 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affv ID | Genbank | Seg ID | Known Gene Name | fold change in metas | r p value | metastati c: Mean | metastati metastati c: c: Std Median Dev | metastati c: Std Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|--|--------------------------------------|--|--|--------------------------------|-----------------------------------|-------------------------------------|--|--------------------------------------|--|--|---|
| S68287_at | S68287 | alc 1, rec hyy de de dib | aldo-keto reductase family 1, member C4 (chlordecone reductase; 3-alpha hydroxysteroid dehydrogenase, type I; dihydrodiol dehydrogenase 4) | 24.11 | 0 | 6.23 | 6.37 | 9.75 | 510.59 | 569.32 | 165.14 |
| S69232_at | S69232 | 3027 f | electron-transferring- 3027 flavoprotein dehydrogenase | 4.9 | 0.00017 | 2.73 | 4.9 | 4.29 | 113.33 | 103.27 | 60.93 |
| S70004_at | S70004 | 3029 | 3029 glycogen synthase 2 (liver) | 96.6 | 0.00001 | -0.03 | -4.28 | 11.42 | 225.1 | 212.32 | 102.76 |
| S72370_at | S72370 | 3030 | 3030 pyruvate carboxylase | 5.31 | 0.00075 | 3.83 | 5.81 | 23.1 | 113.03 | 118.49 | 50.19 |
| S77410_at | S77410 | 3035 | angiotensin receptor 1 ras homolog gene family. | 10.4 | 0 | 18.77 | 17.34 | 10.85 | 266.59 | 284.37 | 98.14 |
| S82240_at | S82240 | | member E | #N/A | #N/A | 19.9 | 6.78 | 30.8 | 75.14 | 74.07 | 40.43 |
| T08879_at rc_T16306_at | T08879 T16306 | 3049 0 | cathepsin F EST | 5.29 #N/A | 0.0008 #N/A | 17.98 24.23 | 14.06 23.64 | 12.92 24.91 | 156.33 58.82 | 180.88 63.62 | 79.67 22.35 |
| rc_T40895_at rc_T46901_at rc_T48075_f_at rc_T50773_s_at | T40895 T46901 T48075 T50773 | | protein tyrosine phosphatase type IVA, member 1 EST hemoglobin, alpha 1 apolipoprotein C-IV | #N/A 77.28 32.56 #N/A | #N/A 0.0006 0.00172 #N/A | 1575.59 11.33 246.19 94.77 | 2039.82 -31.1 164.89 43.61 | 1098.29 121.66 271.65 169.6 | 4148.86 2608.54 7193.85 8473.97 | 2440.28 2853.98 5199.28 6484.47 | 4169.52 691.43 6317.01 7036.85 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|---|----------------------------|----------------------|---|------------------------|-------------------------|------------------------|--------------------------|-------------------------|-----------------------------|-----------------------------|----------------------------|
| Affy ID | Genbank | Seq ID | Known Gene Name | change in metas | p value | metastati c: Mean | c: Median | c: Std Dev | set 2: Mean | set 2: Median | set 2: Std Dev |
| rc_T51617_at rc_T51930_at | T51617 T51930 | 3138 3139 | solute carrier family 22 (extraneuronal monoamine transporter), member 3 EST | 17.67 | 0.00018 0.00066 | 29.08 8.27 | 19.44 9.41 | 56.17 5.97 | 669.34 325.48 | 725.25 218.59 | 402.58 298.12 |
| rc_T56281_f_at T57140_s_at rc_T58756_at | T56281 T57140 T58756 | 3151 3152 3156 | RNA helicase-related protein paraoxonase 3 EST | 32.34 28.8 13.39 | 0.00002 0 0.00013 | 71.16 -1.37 7.38 | 62.87 -7.55 -19.91 | 44.81 22.26 46.26 | 2132.07 825.16 390.04 | 2146.52 979.66 386.76 | 824.76 487.32 164.41 |
| rc_T58775_at | 158775 | 3157 | small inducible cytokine subfamily A (Cys-Cys), member 16 | 21.18 | 0.00006 | 11.93 | -0.59 | 34.83 | 577.14 | 540.96 | 205.27 |
| rc_T59148_s_at | T59148 | 3158 | carbamoyl-phosphate synthetase 1, mitochondrial ketohexokinase | 88.89 | 0 | 17.1 | 2.26 | 49.29 | 3384.09 | 3845.28 | 2295.81 |
| rc_T61256_s_at | T61256 | 3162 | (fructokinase) | 13.59 | 0.00425 | 55.8 | 29.78 | 120.62 | 752.99 | 846.1 | 315.74 |
| rc_161654_r_at rc_T64933_r_at | 161654 T64933 | 3166 | apolipoprotein A-I EST | 4 X # # | #\\# #\\A | 66.98 1.58 | -33.89 10.35 | 208.52 | 9388.4 52.88 | 4297.02 40.46 | 12058.3 54.68 |
| rc_T65443_s_at | T65443 | 3175 | EST | #N/A | #N/A | 10.45 | 9.99 | 13.16 | 70.03 | 75.01 | 29 |
| rc_T66189_s_at rc_T67520_at | T66189 T67520 | 3178 3182 | glutaryl-Coenzyme A dehydrogenase EST | 8.61 #N/A | 0.00003 #N/A | 17.08 52.46 | 20.49 49.25 | 23.82 28.02 | 269.19 105.56 | 301.13 113.95 | 136.96 38.23 |
| rc_T67705_s_at | T67705 | 3183 | 3183 asialoglycoprotein receptor 2 | 31.6 | 0.00705 | 168.64 | 37.91 | 284.26 | 2403.76 | 2437.4 | 683.22 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|------------------------|------------------|----------------|--|----------|---------|-----------|---------------------|-----------|---------|---------|---------|
| | | | | change | | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank Seq ID | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc_T67931_at | T67931 | fil 3184 pt | fibrinogen, B beta 3184 polypeptide | 49.55 | 0 | -22.74 | -22.4 | 23.54 | 1187.95 | 1000.67 | 692.18 |
| | i co | S | short-chain | 2 | 0.0450 | , , | 7 | 70 73 | 262 22 | 200.22 | 126 49 |
| 168083_at T68510_at | 168083 T68510 | 3185 Q | 3185 denydrogenase/reductase 1 3187 EST | 3.87 | 0.00617 | 0.88 | 47.39 -1.97 | 11.33 | 113.31 | 94.26 | 103.92 |
| rc T68711 at | T68711 | 3188 E | ST | 37.65 | 0.00036 | 19.73 | -26.69 | 95.05 | 1388.67 | 1074.67 | 890.91 |
| rc_T68855_at | T68855 | 3189 E | EST | 34.04 | 0 | 8.38 | 6.98 | 6 | 800.27 | 911.26 | 406.27 |
| rc T69020 s at | T69020 | 3193 E | EST | 5.39 | 0.00383 | 32.69 | 17.15 | 46.31 | 202.05 | 208.64 | 90.05 |
| rc_T69164_at | T69164 | 3195 E | EST | 4.38 | 0.00548 | 34.31 | 17.81 | . 30.85 | 173.2 | 167.2 | 85.34 |
| | | E & | mannose-binding lectin (protein C) 2, soluble | | | | | | | | |
| rc T69284 s at | T69284 | 3196 (c | psonic defect) | 36.53 | 0 | 9.82 | 0.32 | 17.14 | 1147.2 | 618.07 | 972.57 |
| rc_T69305_at | T69305 | 3197 E | 3197 EST | #N/A | #N/A | -107.09 | -119.3 | 45.51 | 4052.6 | 4262.37 | 3025.26 |
| T69384 at | T69384 | | period (Drosophila) nomolog 1 | 5.88 | 0.01219 | 37.94 | 19.8 | 90.03 | 232.84 | 205.78 | 126.44 |
| rc_T71978_at | T71978 | 3205 E | EST | 4.39 | 0.0017 | 41.46 | 58.86 | 43.93 | 218.38 | 208.73 | 89.39 |
| rc T72171 s at | T72171 | 3206 th | lyroxin-binding globulin | 10.41 | 0.00163 | 5.48 | -20.52 | 54.58 | 492.89 | 371.71 | 454.89 |
| rc T72502 at | T72502 | 3208 E | 3208 EST | 8.87 | 0.0000 | 24.11 | 24.41 | 28.19 | 308.87 | 271.27 | 179.15 |
| rc_T72906_at | T72906 | 3210 E | ST | 7.56 | 0.00062 | 4.88 | 3.04 | 14.04 | 233.73 | 199.26 | 211.99 |
| rc_T73442_at | T73442 | 3213 E | ST | 94.41 | 0 | 28.55 | 25.96 | 25.95 | 3170.29 | 3275.73 | 1073.74 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | 2 | | fold change | | metastati | metastati metastati c: c: Std | metastati c: Std | normal set 2: | normal set 2: | normal set 2: |
|---|--|--|--|---|---|---|---|--|---|--|--|
| Ally ID | Gendank Seq ID | 71 bac | Allowii Gene Name | In metas | p value | C: Mean | Median | Dev | Mean | Median | Sta Dev |
| rc_T73739_s_at | T73739 | 3214 | alanine-glyoxylate aminotransferase (oxalosis I; hyperoxaluria I; glycolicaciduria; serine- pyruvate aminotransferase) | 16.18 | 0.00277 | 1891.22 | 896.45 | 2229.6 | 26094.38 | 15281.08 | 25947.4 |
| rc_T74542_s_at | T74542 | 3215 | UDP glycosyltransferase 2 3215 family, polypeptide B10 | 32.36 | 0.00004 | 27.96 | 25.04 | 64.23 | 1374.78 | 1445.94 | 869.3 |
| rc_T74608_at | T74608 | . 3216 | hydroxyacid oxidase . 3216 (glycolate oxidase) 1 | 14.03 | 0 | -3.14 | -1.78 | 23.12 | 360.97 | 328.7 | 194.85 |
| rc_T82323_at | T82323 | 3231 | immunoglobulin superfamily, member 4 | 10.14 | O | 14.47 | 5.07 | 19.91 | 264.33 | 229.92 | 135.46 |
| T83397_at rc_T84084_at | T83397 T84084 | 3233 3234 | phytanoyl-CoA hydroxylase (Refsum disease) EST | 63.6 #N/A | 0 #N/A | -17.97 21.47 | -4.64 21.35 | 35.51 9.96 | 1404.6 68.96 | 1421.69 66.69 | 605.99 27.4 |
| rc_T84491_at rc_T86482_at rc_T87224_at rc_T90037_at rc_T90520_at rc_T91161_at | T84491 T86482 T87224 T90037 T90520 | 3235 3238 3241 3246 3249 3253 | CUG triplet repeat,RNA- binding protein 2 transferrin EST EST EST | #N/A 11.15 #N/A 3.99 8.67 3.48 | #N/A 0.00006 #N/A 0.0016 0.00072 0.00002 | 32.84 7.6 584.24 23.17 9.05 9.95 | 34.93 9.41 451.54 25.82 -7.07 | 8.15 12.34 184.49 23.62 25.65 11.55 | 79.08 316.82 909.33 150.67 306.39 | 80.41 197.03 851.86 107.18 362.5 70.8 | 33.03 302.36 431.47 126.19 204.34 30.28 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | netastati | normal | normal | normal |
|--------------------------|---------|--------------|--|---------------|---------|----------------|---------------------|---------------|-------------------|------------------|------------------|
| | | | | change | | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | 0, | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| T95813_f_at | T95813 | | 3262 KIAA1051 protein | 38.38 | 0.00008 | 56.83 | 42.22 | 56.86 | 2271.29 | 1809.29 | 1537.82 |
| U00115_at | U00115 | 3272 | B-cell CLL/lymphoma 6 (zinc finger protein 51) | 5.18 | 0.00045 | 16.17 | 10.31 | 26.94 | 132:13 | 133.88 | 32.06 |
| U01120_at | U01120 | 3274 | glucose-6-phosphatase, catalytic (glycogen storage disease type I, von Gierke 3274 disease) | 13.41 | 0.00147 | 83.05 | 45.5 | 91.8 | 878.33 | 693.43 | 495.6 |
| U02388_at | U02388 | 3278 | cytochrome P450, subfamily IVF, polypeptide 2 | 9.38 | 0.00001 | -4.93 | -3.43 | 18.14 | 207.48 | 207.15 | 90.12 |
| U03056_at | U03056 | 3280 | 3280 hyaluronoglucosaminidase 1 | 6.64 | 0 | 12.68 | 11.77 | 4.73 | 139.32 | 137.21 | 43.97 |
| U05861_at | U05861 | 3285 | aldo-keto reductase family 1, member C1 (dihydrodiol dehydrogenase 1; 20-alpha (3-alpha)-hydroxysteroid dehydrogenase) | 23.22 | 0.00002 | 47.98 | 44.41 | 21.91 | 1070.69 | 962.74 | 359.01 |
| U06641_s_at U06863_at | U06863 | 3287 3288 | UDP glycosyltransferase 2 3287 family, polypeptide B15 3288 follistatin-like 1 | 24.32 3.09 | 0.00001 | 56.39 43.48 | 62.92 | 54.9 14.83 | 1762.46 141.97 | 1615.25 149.4 | 1206.06 59.45 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold change | | metastati | metastati metastati c: c: Std | metastati c: Std | normal set 2: | normal set 2: | normal set 2: |
|------------------------|------------------|--------------|---|----------------|--------------|----------------|----------------------------------|---------------------|------------------|------------------|------------------|
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| U08006_s_at | | 3290 | complement component 8, alpha polypeptide | 9 | 0.00111 | 47.06 | 31.92 | 47.52 | 633.01 | 661.72 | 102.98 |
| U08198_rna1_at | 008198 | 3292 | complement component 8, gamma polypeptide | 19.71 | 0 | -29.27 | -38.32 | 47.54 | 544.28 | 630.51 | 234.87 |
| U08854_s_at | U08854 | 3293 | UDP glycosyltransferase 2 3293 family, polypeptide B15 | 7.38 | 0.00005 | 10.72 | 6.45 | 10.71 | 181.08 | 184.97 | 91.81 |
| U11313_at | U11313 | 3297 | 3297 sterol carrier protein 2 | 5.44 | 0.00187 | 17.39 | 0.82 | 35.86 | 181.8 | 228.15 | 99.25 |
| U12778_at | U12778 | 3301 | acyl-Coenzyme A dehydrogenase, short/branched chain | 5.68 | 0.00116 | 5.23 | 6.16 | 7.33 | 147.06 | 158.14 | 91.5 |
| U13061_ma1_at | U13061 | 3302 | sulfotransferase family 2A, dehydroepiandrosterone (DHEA) -preferring, member | 11.32 | 0.00048 | 98.6 | 7.49 | 19.46 | 369.12 | 431.45 | 213.83 |
| U15174_at | U15174 | 3310 | BCL2/adenovirus E1B 19kD-3310 interacting protein 3 | #N/A | #N/A | 41.72 | 11.86 | 49.01 | 129.19 | 138.02 | 68.72 |
| U15932_at U17989_at | U15932 U17989 | 3311 3316 | dual specificity phosphatase 5 nuclear autoantigen | #N/A #N/A | #N/A #N/A | 47.94 14.21 | 27.05 12.72 | 47.87 6.13 | 86.73 44.22 | 86.97 | 38.45 19.09 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| ı | | | | | fold change | | metastati | metastati metastati c: c: Std | netastati c: Std | normal set 2: | normal set 2: | normal set 2: |
|---|------------------------|------------------|------------------------|--|----------------|--------------|-----------------|----------------------------------|---------------------|------------------|------------------|------------------|
| l | Affy ID | Genbank Seq ID | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| | U19495_s_at | U19495 | 3319 stı | 3319 stromal cell-derived factor 1 | 4.59 | 0.00011 | 7.17 | 5.76 | 5.65 | 103.81 | 114.61 | 50.36 |
| | U19523_at | U19523 | G 3320 re | GTP cyclohydrolase 1 (dopa- 3320 responsive dystonia) | 5.25 | 0.00029 | 12.84 | 12.26 | 15.75 | 136.86 | 142.71 | 66.64 |
| | U20530_at | U20530 | se 3322 24 | secreted phosphoprotein 2, 24kD | 16.93 | 0 | 12.63 | 11.54 | 9.93 | 383.92 | 421.24 | 128.02 |
| | U20938_at | U20938 | 3324 de | uiriya opyiiiniai e 3324 dehydrogenase | 3.92 | 0.00053 | 99.9 | 90.9 | 7.12 | 99.06 | 97.02 | 44.64 |
| | U22029_f_at | U22029 | cy \ 3327 pc | cytochrome P450, subfamily IIA (phenobarbital-inducible), 3327 polypeptide 7 | 71.98 | 0 | 5.05 | 10.11 | 21.51 | 1968.7 | 1339.04 | 1937.45 |
| _ | U22662_at U31342_at | U22662 U31342 | 3329 1, 3344 nu | nuclear receptor subfamily 1, group H, member 3 nucleobindin 1 | #N/A #N/A | #N/A #N/A | 5.71 | 14.46 | 20.18 40.03 | 63.21 183.59 | 58.72 | 37.5 58.86 |
| _ | U32576_rna1_at | U32576 | | apolipoprotein C-IV | 18.59 | 0.00005 | 511.9/ -7.01 | 516.75 -23.2 | 62.7 | 606.88 | 608.66 | 224.72 |
| | U32989_at | U32989 | 3348 try | 3348 tryptophan 2,3-dioxygenase | 10.07 | 0.02825 | 166.12 | 22.7 | 333.99 | 527.61 | 473.11 | 268.8 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affy ID | Genbank | Seq ID | Known Gene Name | fold change in metas | p value | metastati c: Mean | metastati metastati c: c: Std Median Dev | metastati c: Std Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|------------------------|------------------|--------------|--|----------------------------|--------------|----------------------|--|----------------------------|--------------------------|----------------------------|-----------------------------|
| U37055_ma1_s_at | U37055 | 3353 | macrophage stimulating 1 (hepatocyte growth factor- like),macrophage stimulating, pseudogene 9 | 18.28 | 0 | -7.37 | -38.2 | 47.48 | 549.49 | 626.51 | 217.69 |
| U37143_at U37547_at | U37143 U37547 | 3354 | cytochrome P450, subfamily IIJ (arachidonic acid epoxygenase) polypeptide 2 apoptosis inhibitor 1 | #N/A #N/A | #N/A #N/A | 33.79 29.52 | 28.24 19.51 | 31.46 48.74 | 116.06 93.6 | 131.46 112.45 | 48.68 52.27 |
| U39226_at | U39226 | 3358 | myosin VIIA (Usher syndrome 1B (autosomal recessive, severe)) | #N/A | #N/A | 33.54 | 31.07 | 8.59 | 53.24 | 50.47 | 20.86 |
| U46689_at | U46689 | 3366 | aldehyde dehydrogenase 10 (fatty aldehyde dehydrogenase) | 3.5 | 0.01292 | 31.22 | 25.36 | 49.19 | 124.41 | 136.16 | 53.23 |
| U48707_at U49082_at | U48707 U49082 | 3371 3373 | protein phosphatase 1, regulatory (inhibitor) subunit 1A transporter protein | 14.22 | 0.00088 | -1.33 | -1.45 46.71 | 20.92 56.13 | 319.97 563.34 | 310.24 515.31 | 125.51 179.82 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affv ID | Genbank Sed ID | Sed ID | Known Gene Name | fold change in metas | o value | metastati c: Mean | metastati metastati c: c: Std Median Dev | metastati c: Std Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|-------------|----------------|--------|---|----------------------------|---------|----------------------|--|----------------------------|--------------------------|----------------------------|-----------------------------|
| U49248_at | U49248 | 3374 | ATP famil men | 7.06 | 0.00067 | 30.01 | 26.03 | 36.14 | 231.96 | 249.25 | 91.2 |
| U50929_at | U50929 | 3380 | | 36.91 | 0 | 8.63 | 2.82 | 20.35 | 898.57 | 865.9 | 313.61 |
| U51010_s_at | U51010 | 3381 | nicotinamide N- methyltransferase | 27.79 | 0.00366 | 105.25 | 43.72 | 204.23 | 1589.66 | 1590.62 | 536.04 |
| U56814_at | U56814 | 3393 | 3393 deoxyribonuclease I-like 3 | 29.43 | 0.00003 | 5.04 | 11.75 | 22.3 | 853.78 | 1073.24 | 434.79 |
| U57094_at | U57094 | 3394 | | #N/A | #N/A | 30.85 | 31.58 | 23.86 | 93.06 | 93.79 | 50.75 |
| U59111_at | U59111 | 3397 | dermatan sulpnate proteoglycan 3 | #N/A | #N/A | 32.34 | 26.73 | 15.11 | 69.89 | 64.44 | 32.48 |
| U59423_at | U59423 | 3399 | MAD (mothers against decapentaplegic, Drosophila) homolog 1 | #N/A | #N/A | 17.46 | 16.81 | 10.17 | 60.5 | 51.09 | 37.25 |
| U60061_at | U60061 | 3400 | fasciculation and elongation 3400 protein zeta 2 (zygin II) | #N/A | #N/A | 61.44 | 29.18 | 68.3 | 55.02 | 66.29 | 38.2 |
| U66672_at | U66672 | 3407 | ATP-binding cassette, sub- 3407 family A (ABC1), member 5 | #N/A | #N/A | 78.03 | 41.24 | 72.76 | 105.45 | 97.64 | 60.15 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | 1 | metastati metastati | metastati | normal | normal | normal |
|------------------------|------------------|------------------|---|--------------------|--------------------|----------------------|---------------------|----------------|----------------|------------------|-------------------|
| Affy ID | Genbank | Seq ID | Known Gene Name | change in metas | p value | metastati c: Mean | c: Median | c: Std Dev | set 2: Mean | set 2: Median | set 2: Std Dev |
| U68233_at | U68233 | 3412 | nuclear receptor subfamily 3412 1, group H, member 4 | 92.9 | 0.00022 | 19.94 | 89 | 29.99 | 198.25 | 233.6 | 83.71 |
| U69141_at | U69141 | 3414 0 | glutaryl-Coenzyme A 3414 dehydrogenase | 3.06 | 0.00053 | 35.13 | 27.91 | 13.89 | 108.36 | 119.26 | 34.82 |
| U70732_rna1_at | U70732 | 3415 E | glutamic-pyruvate transaminase (alanine aminotransferase) | 27.63 | 0 | -47.95 | -65.21 | 50.88 | 696.22 | 732.8 | 284.7 |
| U70732_rna1_at | U70732 | 9 t 3415 | glutamic-pyruvate transaminase (alanine aminotransferase) | 27.63 | . 0 | -1.44 | 3.98 | 66.42 | 201.88 | 218.68 | 136.17 |
| U72515_at | U72515 | 3416 r | putative protein similar to nessy (Drosophila) | #N/A | #N/A | 45.44 | 58.93 | 45.12 | 137.35 | 138.01 | 55.38 |
| U73514_at | U73514 | 3419 | hydroxyacyl-Coenzyme A 3419 dehydrogenase, type II | 3.79 | 0.01392 | 9.85 | -22.7 | 56.91 | 159.74 | 161.9 | 127.8 |
| U77396_at | U77396 | 1 3426 f | LPS-induced TNF-alpha factor | 4.47 | 0.00108 | 30.53 | 33.74 | 31.54 | 169.83 | 141.36 | 87.43 |
| U77594_at U79716_at | U77594 U79716 | 3428 i 3437 r | retinoic acid receptor responder (tazarotene induced) 2 | 11.95 3.51 | 0.00527 0.00053 | 94.59 14.44 | 122.86 9.15 | 71.01 15.58 | 838.98 90.6 | 864.02 101.49 | 313.87 36.51 |
| U80226_s_at | U80226 | 3439 8 | 4-aminobutyrate aminotransferase | #N/A | #N/A | 39.69 | 51.46 | 20.33 | 86.94 | 91.76 | 24.15 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| I | Affv ID | Genbank | Seq ID | Known Gene Name | fold change in metas | p value | metastati c: Mean | metastati metastati c: c: Std Median Dev | metastati c: Std Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|----------|---|--------------------------------------|------------------------------|--|------------------------------|-----------------------------------|-------------------------------|--|----------------------------------|------------------------------------|------------------------------------|-----------------------------------|
| l | U81607_at | U81607 | 3440 | A kinase (PRKA) anchor protein (gravin) 12 solute carrier family 9 | #N/A | #N/A | 18.75 | 8.46 | 50.91 | 82.7 | 78.31 | 62.79 |
| | U82108_s_at | U82108 | 3441 | (sodium/hydrogen exchanger), isoform 3 regulatory factor 2 | 3.19 | 0.01545 | 58.61 | 78.15 | 43.49 | 130.34 | 122.39 | 73.74 |
| | U83461_at U85193_at | U83461 U85193 | 3444 3447 | solute carrier family 31 (copper transporters), member 2 nuclear factor I/B | #N/A #N/A | #N/A #N/A | 24.64 21.84 | 21.28 | 13.16 | 51.79 72.71 | 48.53 68.69 | 20.85 32.02 |
| | U90544_at | U90544 | 3454 | solute carrier family 17 (sodium phosphate), member 2 | 6.42 | 0.00023 | -3.14 | 5.18 | 19.12 | 157.41 | 168.92 | 63.8 |
| | U95090_at | 06056N | 3464 | nephrosis 1, congenital, Finnish type (nephrin) | 14.01 | 0.00018 | 40.49 | 30.15 | 38.71 | 554.93 | 539.41 | 183.55 |
| | U95090_at W02027_s_at W03796_at rc_W15417_at | U95090 W02027 W03796 W15417 | 3464 3465 3468 3473 | nephrosis 1, congenital, Finnish type (nephrin) EST EST KIAA0699 protein | 14.01 #N/A 8.3 #N/A | 0.00018 #N/A 0.0032 #N/A | 4.97 7.28 8.13 30.82 | 5.87 2.64 -19.16 20.85 | 18.32 14.09 75.11 38.03 | 188.83 30.56 336.51 83.24 | 201.08 28.52 383.56 81.29 | 88.01 12.47 190.14 49.83 |
| | W16686_at | W16686 | 3476 | basic helix-loop-helix domain containing, class B, 2 | #N/A | #N/A | 46.29 | -6.45 | 123.94 | 100.9 | 83.13 | 79.01 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | norma |
|---------------|----------|----------|--|----------|----------|-----------|---------------------|-----------|---------|---------|---------|
| | | | | change | | metastati | ၓ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| W27503 at | W27503 | 3486 KIA | KIAA0679 protein | #N/A | #N/A | 17.31 | 17.31 | 21.95 | 27.71 | 27.51 | 5.28 |
| W28235_at | W28235 | 3487 | EST | #N/A | #N/A | 95.97 | 150.37 | 81.4 | 271.36 | 307.26 | 108.42 |
| W28824_at | W28824 | 3493 | EST | 15.8 | 0.00006 | -1.99 | -5.59 | 17.64 | 441.67 | 509.29 | 251 |
| | | | Kreisler (mouse) maf-related | | | | | | | | |
| 14/26200 6 64 | 14/26200 | 2504 | la iona shoot homolog | 700 | 0.02748 | 40 52 | 70 4 | 72.24 | 1577 | 156 24 | 77 72 |
| W3023U_S_at | 06206 VV | | reactifie zipper fromotog | 9.0 | 0.027.10 | 40.02 | 10.43 | 13.61 | 1.701 | 130.24 | 03.13 |
| W38//8_s_at | W38//8 | | ESI | #N/A | #N/A | -55.97 | -63.87 | 44.35 | /8.UJ | 37.07 | 137.01 |
| rc_W42429_at | W42429 | 3209 | EST | #N/A | #N/A | 98.32 | 80.07 | 52.23 | 252.63 | 300.68 | 162.35 |
| rc_W45560_at | W45560 | 3525 | EST | 5.06 | 0.00996 | 86.81 | 62.95 | 81.92 | 369.61 | 354.89 | 158.02 |
| rc_W48860_at | W48860 | 3537 | EST | #N/A | #N/A | 36.89 | 33 | 14.08 | 52.69 | 41.31 | 21.1 |
| | | | CD36 antigen (collagen type I receptor, thrombospondin receptor)-like 2 (lysosomal integral membrane protein | | • | | | | | | |
| W52581_at | W52581 | 3543 | î (| 3.08 | 0.00091 | 10.92 | 8.89 | 10.63 | 71.35 | 58.02 | 37.28 |
| W58520 at | W58520 | 3552 | nucleoporin 88kD | #N/A | #N/A | 24.77 | 26.75 | 16.56 | 81.88 | 87.47 | 37.77 |
| rc W58756 at | W58756 | 3554 | EST | 12.17 | 0.00087 | 93.37 | 88.97 | 71.48 | 1363.71 | 1064.26 | 1032.27 |
| rc_W60186_at | W60186 | | EST | 3.24 | 0.00228 | 19.15 | 26.5 | 27.35 | 101.92 | 109.68 | 47.85 |
| rc_W63741_at | W63741 | 3566 | EST | #N/A | #N/A | -11.85 | -13.73 | 14.25 | 28.54 | 17.1 | 44.58 |
| rc_W67147_at | W67147 | 3568 | deleted in liver cancer 1 | 9.74 | 0.00002 | 16.09 | 9.11 | 24.66 | 282.59 | 235.29 | 169.33 |
| rc_W67199_at | W67199 | 3569 | EST | 3.09 | 0.01528 | 30.05 | 35 | 12.16 | 123.27 | 117.99 | 90.47 |
| rc_W70131_at | W70131 | 3577 | EST | #N/A | #N/A | 108.21 | 30.68 | 183.05 | 118.49 | 100.66 | 53.4 |
| rc_W72044_at | W72044 | | insulin induced gene 1 | 24.58 | 0.00001 | 41.6 | 36.28 | 36.46 | 1133.09 | 940.16 | 723.6 |
| rc_W72079_at | W72079 | 3581 | EST | 6.36 | 0.00641 | 46.78 | 17.94 | 66.71 | 261.7 | 214.62 | 175.87 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|--------------------------------|------------------|--------------|--|---------------|-----------|--------------|---------------------|---------------|------------------|------------------|-----------------|
| | | | | change | | metastati | ຮ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank Seq ID | Seq ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean . | Median | Std Dev |
| | | | oxidative 3 alpha | | | | | | , | | |
| | | | hydroxysteroid dehydrogenase; retinol | | | | | | | | |
| rc W72382 at | W72382 | 3584 | dehydrogenase | #N/A | #N/A | 37.21 | 6.21 | 66.91 | 3403.46 | 3255.54 | 1538.23 |
| rc_W72471_at | W72471 | 3585 | | #N/A | #N/A | 17.93 | 13.32 | 10.61 | 71.51 | 76.9 | 29.33 |
| rc_W72972_at | W72972 | 3587 | | 5.61 | 0.00939 | 44.7 | 26.44 | 80.59 | 277.4 | 270.63 | 183.07 |
| rc_W73382_at | W73382 | 3591 | EST | #N/A | #N/A | 62.97 | 39.05 | 67.26 | 87.32 | 89.49 | 28.17 |
| rc W79422 s at | W79422 | 3604 | fumarylacetoacetate | 14.94 | 0.00059 | -18.4 | -19.27 | 33.52 | 467.37 | 554.19 | 306.71 |
| W81053 at | W81053 | 3610 | EST | 4.91 | 0.00164 | -21.38 | -13.04 | 24.17 | 135.84 | 109.34 | 113.36 |
| rc W85765 at | W85765 | 3618 | EST | 3.83 | 0.00379 | 3.6 | 1.46 | 7.87 | 108.99 | 77.01 | 109.58 |
| rc_W85847_at | W85847 | 3619 | EST | 7.28 | 0.00024 | 4.15 | 5.29 | 9.26 | 187.31 | 160.02 | 130.73 |
| rc_W85886_at | W85886 | 3621 | EST | 3.46 | 0.00814 | -6.09 | -2.4 | 11.46 | 99.92 | 68.44 | 97.4 |
| l | | • | protein C inhibitor (plasminogen activator | | | | | | | | |
| rc W86431 s at | W86431 | 3627 | inhibitor III) | 14.94 | 0.00152 | 3.16 | 5.52 | 12.03 | 606.44 | 395.34 | 590.7 |
| rc_W86600_at | W86600 | 3628 | EST | 32.14 | 0 | 12.67 | 7.49 | 15.61 | 792.37 | 807.74 | 313.83 |
| | | | homogentisate 1,2- | | | | | | | | |
| | | | dioxygenase (homogentisate | | | | | | | | |
| rc_W87454_at | W87454 | 3632 | oxidase) STAT induced STAT | 7.93 | 0.00149 | 11.36 | 5.83 | 39.02 | 336.07 | 275.03 | 315.24 |
| rc_W87480_at | W87480 | 3633 | inhibitor-2 protein Z, vitamin K- | 3.24 | 0.01063 | 31.91 | 26.07 | 30.44 | 129.06 | 119.73 | 80.94 |
| rc_W87606_s_at | W87606 | 3635 | | 9.23 | 0.00085 | 2.53 | -1.74 | 96.6 | 268.59 | 280.44 | 184.7 |
| rc_W88946_at rc_W90018_f_at | W88946 W90018 | 3639 3642 | | 58.26 #N/A | o #N/A | 2.16 -2.1 | -22.13 4.12 | 39.9 13.16 | 1717.54 59.32 | 1821.77 53.19 | 502.55 41.68 |
| | | | | | | | | | | | |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|----------------|----------------|----------|---|----------|---------|-----------|---------------------|-----------|---------|---------|---------|
| | | • | | change | _ | metastati | ដ | c: Std | set 2: | set 2: | set 2: |
| Affy ID | Genbank Seg ID | Sed ID | Known Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Std Dev |
| rc_W90396_at | W90396 | 3645 EST | EST | #N/A | #N/A | 16.1 | 15.26 | 11.11 | 23.65 | 22.93 | 10.28 |
| rc_W92148_s_at | W92148 | 3650 | 3650 kininogen | 51.09 | 0.00376 | 220.31 | 42.96 | 427.59 | 3697.54 | 4015.04 | 1444.99 |
| rc_W94942_i_at | W94942 | 3661 | dadi openining proopriated 10 | 7.23 | 0.00137 | 19.36 | 16.05 | 11.9 | 250.16 | 281.49 | 160.43 |
| rc_W95041_at | W95041 | 3662 | EST | 12.79 | 0.00001 | 99.9 | 5.17 | 3.55 | 291.2 | 298.05 | 115.81 |
| X02160_at | X02160 | 3671 | insulin receptor | 5.29 | 0.0001 | 10.06 | 8.21 | 10.89 | 77.32 | 65.71 | 45.53 |
| X02176_s_at | X02176 | 3672 | 3672 complement component 9 protein C (inactivator of | 9.61 | 0.00004 | 47.28 | 46.34 | 28.54 | 507.56 | 507.21 | 261.45 |
| X02750_at | X02750 | 3673 | coagulation factors Va and VIIIa) | 7.04 | 0.00079 | 55.79 | 52.95 | 30.08 | 373.11 | 367.46 | 95.74 |
| X06562_at | X06562 | 3686 | 3686 growth hormone receptor | 10.87 | 0.00001 | 10.46 | 68.9 | 15.53 | 291.31 | 314.9 | 160.29 |
| X07618_s_at | X07618 | 3691 | cytochrome P450, subfamily IID (debrisoquine, sparteine, etc., -metabolising), polypeptide 7a (pseudogene) | 35.79 | 0.00065 | 59.61 | 12.32 | 101.97 | 1429.98 | 1631.04 | 671.55 |
| X07618_s_at | X07618 | 3691 | cytochrome P450, subfamily IID (debrisoquine, sparteine, etc., -metabolising), polypeptide 7a 3691 (pseudogene) | 35.79 | 0.00065 | 40.81 | 38.21 | 14.35 | 609.22 | 605.91 | 330.02 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affy ID | Genbank Seq ID | Seq ID | Known Gene Name | fold change in metas | p value | metastati c: Mean | metastati metastati c: c: Std Median Dev | metastati c: Std Dev | normal set 2: Mean | normal set 2: Median | normal set 2: Std Dev |
|-----------------------------|------------------|--------------|--|----------------------------|----------------|----------------------|--|----------------------------|--------------------------|----------------------------|-----------------------------|
| X07619_s_at | X07619 | 3692 | cytochrome P450, subfamily IID (debrisoquine, sparteine, etc., -metabolising), polypeptide 7a (pseudogene) | 4.96 | 0.00004 | 32.26 | 33.56 | 10.88 | 177.56 | 206.15 | 77.69 |
| X07732_at | X07732 | 3693 | hepsin (transmembrane protease, serine 1) | 28.21 | 0 | -8.34 | -12 | 39.24 | 678.15 | 672.42 | 183.61 |
| X07767_at X12662_ma1_at | X07767 X12662 | 3694 3697 | protein kinase, cAMP- dependent, catalytic, alpha arginase, liver | 5.94 20.59 | 0.00028 | 16.15 11.33 | 19.43 10.15 | 11.48 | 152.58 486.71 | 160.88 479.11 | 73.73 199.71 |
| X13227_at | X13227 | 3698 | 3698 D-amino-acid oxidase | 4.81 | 0.0002 | 25.1 | 33.95 | 26.2 | 166.22 | 147.21 | 77.49 |
| X13930_f_at X14787_at | X13930 X14787 | 3700 3704 | cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 6 thrombospondin 1 | 38.52 3.69 | 0.01115 | 52.66 21.81 | 64.02 | 31.42 | 2408.66 | 1948.18 86.81 | 1843.46 166.32 |
| X14813_at X15393_rna1_at | X14813 X15393 | 3705 3707 | acetyl-Coenzyme A acyltransferase 1 (peroxisomal 3-oxoacyl- Coenzyme A thiolase) motilin | 11.61 #N/A | 0.0002 #N/A | 59.4 70.16 | 71.51 55.83 | 42.45 27.07 | 712.58 113.93 | 704.45 | 258.91 26.14 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | netastati | normal | normal | normal |
|-------------|---------|--------|--|--------------------|---------|----------------------|---------------------|---------------|----------------|------------------|-------------------|
| Affy ID | Genbank | Ol pes | Known Gene Name | change In metas | p value | metastatí c: Mean | c: Median | c: Std Dev | set 2: Mean | set 2: Median | set 2: Std Dev |
| X15422_at | X15422 | 3708 | mannose-binding lectin (protein C) 2, soluble (opsonic defect) | 8.29 | 0.00015 | 2.28 | 2.69 | 7.25 | 208.25 | 221.14 | 123.22 |
| X16260_s_at | X16260 | 3710 | inter-alpha (globulin) inhibitor, H1 polypeptide hepatocyte growth factor | 44.65 | 0.00933 | 40.4 | 36.54 | 29 | 1261.99 | 1133 | 523.31 |
| X16323_at | X16323 | 3711 | | #N/A | #N/A | 10.65 | 6.56 | 6.59 | 32.58 | 30.52 | 9.66 |
| X16663_at | X16663 | 3714 | hematopoletic cell-specific 3714 Lyn substrate 1 | #N/A | #N/A | 48.32 | 28.69 | 56.44 | 81.51 | 83.53 | 33.75 |
| X17094_at | X17094 | 3717 | paired basic amino acid cleaving enzyme (furin, membrane associated receptor protein) | 8.5 | 0 | -1.85 | 7.01 | 25.41 | 190.11 | 216.35 | , 74.11 |
| X53414_at | X53414 | 3728 | alanine-glyoxylate aminotransferase (oxalosis I; hyperoxaluria I; glycolicaciduria; serine- pyruvate aminotransferase) | 13.87 | 0.00003 | 138.92 | 110.29 | | 1856.57 | 1888.79 | 846.06 |
| X54380_at | X54380 | 3730 | 3730 pregnancy-zone protein | 8.44 | 0.00059 | 4.93 | 3.19 | 19.26 | 274.41 | 255.4 | 203.47 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | normal | normal | normal |
|--------------------------|------------------|--------------|---|--------------------|---------|----------------------|---------------------|----------------|------------------|------------------|-------------------|
| Affy ID | Genbank | Seq ID | Known Gene Name | change in metas | p value | metastati c: Mean | c: Median | c: Std Dev | set 2: Mean | set 2: Median | set 2: Std Dev |
| | | | | | | | | | | | |
| X56411_ma1_at | X56411 | 3737 | alcohol dehydrogenase 4 3737 (class II), pi polypeptide | 25.14 | 0.00144 | 55.53 | 11.29 | 102.45 | 991.98 | 943.64 | 448.31 |
| X57025_at X60673_s_at | X57025 X60673 | 3742 3752 | insulin-like growth factor 1 (somatomedin C) adenylate kinase 3 | 4.72 8.3 | 0.00087 | 5.88 23.02 | 10.95 32.17 | 10.22 44.24 | 120.82 331.58 | 100.54 324.75 | 85.16 164.03 |
| 7 01000 | 2000 | , 1 | UDP glycosyltransferase 2 | 9 | | 1 | ! | | | ((| ! |
| A03339_at | Yossaa | 3/28 | ramily, polypeptide B10 c-mvc promofer-binding | 13.00 | 0.00051 | 22.28 | 48.5/ | 41.25 | 669.93 | 732.6 | 161.42 |
| X63417_at | X63417 | 3760 | | #N/A | #N/A | 28.85 | 27.75 | 24.56 | 68.08 | 56.34 | 33.18 |
| X64877_at | X64877 | 3766 | 3766 H factor (complement)-like 3 | 14.6 | 0 | 14.6 | 7.51 | 21.01 | 387.8 | 402.95 | 151.23 |
| X64877_s_at | X64877 | 3766 | 3766 H factor (complement)-like 3 | 12.33 | 0 | 6.14 | 8.26 | 14.25 | 262.65 | 268.29 | 84.84 |
| X65727_cds2_s_at | X65727 | 3768 | glutathione S-transferase A2,glutathione S-transferase A3 | 73.64 | 0 | 21.03 | 17.39 | 15.18 | 2162.22 | 2183.18 | 1435.52 |
| X65962_s_at | X65962 | 3769 | cytochrome P450, subfamily IIC (mephenytoin 4- hydroxylase) | 5.47 | 0.00014 | 36.73 | 31.48 | 11.66 | 247.79 | 149.39 | 208.59 |
| X76105_at | X76105 | 3794 | 3794 death-associated protein . | 3.02 | 0.00944 | 30.12 | 33.72 | 31.36 | 108.68 | 118.36 | 56.75 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati metastati | metastati | norma | normal | normal |
|---|--------------------------------------|------------------------------|--|----------------------|-------------------------|----------------------------------|--------------------------------|---------------------------------|-----------------------------------|--------------------------|-------------------------|
| 4 | 1 | | | change | 1 | metastati | : : | c: Std | set 2: | set 2: | set 2: |
| Arry ID | Gendarik | Sed ID | Anowi Gene Name | III liletas | b value | C. IMEAN | Median | Aan Des | Medil | Median | ara me |
| X78992_at | X78992 | buty 3802 (EGI | butyrate response factor 2 (EGF-response factor 2) | 4.69 | 0.01995 | 64.86 | 31.76 | 66.31 | 318.26 | 336.95 | 207.43 |
| X79981_at | X79981 | 3806 | cadherin 5, VE-cadherin 3806 (vascular epithelium) | #N/A | #N/A | 45.01 | 41.07 | 11.65 | 80.92 | 80.1 | 8.57 |
| X86401_s_at | X86401 | 3815 | glycine amidinotransferase (L-arginine:glycine amidinotransferase) | 22.7 | 0 | 19.17 | 9.16 | 19.13 | 646 | 721.34 | 274.64 |
| X90999_at | 66606X | 3820 | hydroxyacyl glutathione hydrolase; glyoxalase 2 | 5.27 | 0.00047 | 16.15 | 26.83 | 26.47 | 170.37 | 179.54 | 76.98 |
| X91148_at X92396_at X92475_at X94563_vnt2_r_at | X91148 X92396 X92475 X94563 | 3821 3823 3824 3824 | microsomal triglyceride transfer protein (large polypeptide, 88kD) synaptobrevin-like 1 ITBA1 gene | 5.91 #N/A #N/A | 0.00045 #N/A #N/A | 15.37 40.87 56.74 76.79 | 3.85 17.73 63.84 66.4 | 31.49 57.17 25.6 57.64 | 173.77 67.49 66.2 274 91 | 189.11 61.23 63.72 | 83.82 31.24 20.36 |
| X95190_at | X95190 | | | 12.81 | 0 | -80.53 | -76.73 | 37.81 | 285.88 | 349.39 | 119.31 |
| X97160_ma1_at | X97160 | 3838 | transcription factor binding 3838 to IGHM enhancer 3 | #N/A | #N/A | 61.31 | 49.54 | 30.41 | 75.26 | 66.36 | 29.46 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| Affv ID | Genhank | Clipas | Known Gene Name | fold change in metas | <u> </u> | metastati c. Mean | metastati c: Madian | metastati metastati c: c: Std Median Dev | normal set 2: Mean | normal set 2: Median | normal set 2: |
|----------------------------|------------------|--------|---|----------------------------|--------------------|----------------------|---------------------------|--|--------------------------|----------------------------|------------------|
| | | | complement factor H related 3,complement factor H- | | | | | | | | |
| X98337_s_at Y00097_s_at | X98337 Y00097 | 3844 | related 4 annexin A6 | 13.5 3.4 | 0.00001 0.00233 | 18.76 19.43 | 22.2 16.39 | 33.33 33.67 | 411.51 105.39 | 375.08 104.75 | 169.37 39.99 |
| Y00317_at | Y00317 | 3845 | UDP glycosyltransferase 2 3845 family, polýpeptide B4 | 18.34 | 0.00001 | -9.82 | -11.91 | 20.96 | 456.09 | 477.28 | 257.07 |
| Y00317_at | Y00317 | 3845 | UDP glycosyltransferase 2 3845 family, polypeptide B4 | 18.34 | 0.00001 | 11.31 | 14.85 | 7.33 | 231.32 | 282.35 | 122.76 |
| Y00339_s_at | Y00339 | 3847 | 3847 carbonic anhydrase II | 6.89 | 0 | 6.23 | -1.39 | 18.22 | 168.76 | 178.44 | 69.52 |
| Y10659_at | Y10659 | 3859 | interieukin 13 receptor, alpha 1 interieulia 13 montos | 4.22 | 0.00061 | 18.27 | 12.85 | 25.15 | 123.23 | 134.47 | 49.24 |
| Y10659_at | Y10659 | 3859 | meneukin 13 receptor, alpha 1 | 4.22 | 0.00061 | 15.82 | 16.3 | 16.79 | 81.75 | 89.09 | 37.01 |
| Y12711_at | Y12711 | 3861 | progesterone binding protein | 14.83 | 0.00285 | 26.37 | 16.12 | 99.79 | 1061.83 | 935.43 | 829.85 |
| Z28339_at | 228339 | 3875 | aldo-keto reductase family 1, member D1 (delta 4-3- ketosteroid-5-beta- reductase) | 24.66 | 0 | 3.67 | 3.48 | 5.1 | 527.72 | 552.17 | 156.66 |
| Z29481_at | Z29481 | 3877 | 3-hydroxyanthranilate 3,4- 3877 dioxygenase | 6.39 | 0.00029 | 546.48 | 440.33 | 293.12 | 2012.12 | 1797.38 | 944.18 |

Table 9B. Genes and ESTs expressed only in normal liver tissue compared to metastatic liver tissue.

| | | | | fold | | | metastati | metastati metastati | normal | normal | norma |
|----------------|---------|-----------|--|----------|---------|-----------|-----------|---------------------|--------|--------|---------|
| 94 | | ב נ | | change | | metastati | ະ ເ | c: Std | set 2: | set 2: | set 2: |
| Ally ID | Genbank | oed ID | Allown Gene Name | in metas | p value | c: Mean | Median | Dev | Mean | Median | Sta Dev |
| | | _ | nuclear receptor subfamily | | | | | · | | | |
| Z30425_at | Z30425 | 3878 | 1, group I, member 3 | 26.64 | 0 | 5.66 | 7.49 | 8.02 | 181.6 | 192.48 | 76.65 |
| rc_Z38161_at | Z38161 | 3884 | EST | 4.38 | 0.0011 | 11.84 | 12.14 | 6.32 | 111.16 | 107.62 | 80.11 |
| rc_Z38192_s_at | Z38192 | 3885 | EST | 3.35 | 0.00184 | -0.28 | 4.75 | 17.86 | 87.12 | 87.23 | 52.58 |
| rc_Z38910_at | Z38910 | 3898 | EST | #N/A | #N/A | . 89.0 | 2.75 | 12.7 | 35.49 | 32.21 | 23.55 |
| rc_Z39394_at | Z39394 | 3904 | EST | #N/A | #N/A | 10.11 | 10.43 | 11.1 | 24.06 | 22.96 | 11.11 |
| rc_z39622_s_at | Z39622 | | EST | 4.4 | 0.00001 | 19.65 | 21.87 | 11.97 | 110.87 | 115.21 | 36.19 |
| rc_Z39833_at | Z39833 | 3913 (| GTP-binding protein | 16.89 | 0.00034 | 35.79 | 41.07 | 84.59 | 957.61 | 1080.1 | 674.72 |
| rc_Z39976_at | Z39976 | 3915 | EST | 5.76 | 0.00012 | 13.09 | 7.15 | 18.64 | 165.45 | 151.05 | 98.9 |
| rc_Z40259_s_at | Z40259 | 3919 | EST | 8.18 | 0.00002 | 14.39 | 8.61 | 11.98 | 218.4 | 223.24 | 122.06 |
| | | | ariadne, Drosophila, | | | | | | | | |
| rc_z41271_s_at | Z41271 | 3930 | homolog of | #N/A | #N/A | 51.96 | 64.49 | 36.28 | 145.09 | 126.18 | 57.57 |
| rc_Z41356_at | Z41356 | 3932 | EST | #N/A | #N/A | 4.46 | 15.94 | 28.44 | 82.7 | 48.14 | 71.73 |
| rc_Z41634_at | Z41634 | 3934 | EST | #N/A | #N/A | 16.4 | 16.07 | က | 51.19 | 48.49 | 33 |
| l | | _ | flavin containing | | | | | | | | |
| Z47553_at | Z47553 | 3939 | monooxygenase 5 | 6.17 | 0.00011 | 4.86 | 1.26 | 8.96 | 141.98 | 165.57 | 59.33 |
| Z48054_at | 248054 | 3941 | peroxisome receptor 1 | #N/A | #N/A | 46.89 | 51.87 | 23.55 | 78.16 | 80.88 | 26.87 |
| | | . | glucokinase (hexokinase 4) | | | | | | | | |
| Z48475_at | Z48475 | | regulatory protein | 13.84 | 0 | -42.61 | -53.52 | 23.46 | 305.23 | 239.52 | 155.98 |
| Z48633_at | Z48633 | 3944 | EST | W/V# | #N/A | -5.51 | -6.39 | 13.15 | 36.84 | 40.63 | 26.33 |
| | | ., | small inducible cytokine | | | | | | | | |
| Z49269_at | Z49269 | 3945 | subramily A (Cys-Cys), member 14 | 18.46 | 0.00001 | 1.88 | -25.63 | 41.94 | 526.13 | 532.29 | 166.67 |
| Z49878_at | Z49878 | 3946 | guanidinoacetate N- methyltransferase | 13.96 | 0.00021 | 44.63 | 45.34 | 41.76 | 615.57 | 626.06 | 196.31 |

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What is claimed is:

- 1. A method of diagnosing liver cancer in a patient, comprising:
- (a) detecting the level of expression in a tissue sample of two or more genes
 from Tables 3-9; wherein differential expression of the genes in Tables 3-9 is indicative of
 liver cancer.
 - 2. A method of detecting the progression of liver cancer in a patient, comprising:
- (a) detecting the level of expression in a tissue sample of two or more genes from Tables 3-9; wherein differential expression of the genes in Tables 3-9 is indicative of liver cancer progression.
 - 3. A method of monitoring the treatment of a patient with liver cancer, comprising:
 - (a) administering a pharmaceutical composition to the patient;
 - (b) preparing a gene expression profile from a cell or tissue sample from the patient; and
 - (c) comparing the patient gene expression profile to a gene expression from a cell population selected from the group consisting of normal liver cells, hepatocellular carcinoma and metastatic liver carcinoma.
 - 4. A method of treating a patient with liver cancer, comprising:
 - (a) administering to the patient a pharmaceutical composition, wherein the composition alters the expression of at least one gene in Tables 3-9;
- 25 (b) preparing a gene expression profile from a cell or tissue sample from the patient comprising tumor cells; and
 - (c) comparing the patient expression profile to a gene expression profile selected from the group consisting of normal liver cells, hepatocellular carcinoma and metastatic liver carcinoma

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5. A method of diagnosing hepatocellular carcinoma in a patient, comprising:

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- (a) detecting the level of expression in a tissue sample of two or more genes from Tables 3-9; wherein differential expression of the genes in Tables 3-9 is indicative of hepatocellular carcinoma.
- 5 6. A method of detecting the progression of hepatocellular carcinoma in a patient, comprising:
 - (a) detecting the level of expression in a tissue sample of two or more genes from Tables 3-9; wherein differential expression of the genes in Tables 3-9 is indicative of hepatocellular carcinoma progression.

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- 7. A method of monitoring the treatment of a patient with hepatocellular carcinoma, comprising:
 - (a) administering a pharmaceutical composition to the patient;
- (b) preparing a gene expression profile from a cell or tissue sample from the patient; and
 - (c) comparing the patient gene expression profile to a gene expression from a cell population comprising normal liver cells or to a gene expression profile from a cell population comprising hepatocellular carcinoma cells or to both.
 - 8. A method of treating a patient with hepatocellular carcinoma, comprising:
 - (a) administering to the patient a pharmaceutical composition, wherein the composition alters the expression of at least one gene in Tables 3-9;
 - (b) preparing a gene expression profile from a cell or tissue sample from the patient comprising hepatocellular carcinoma cells; and
- 25 (c) comparing the patient expression profile to a gene expression profile from an untreated cell population comprising hepatocellular cells.

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- 9. A method of diagnosing a metastatic liver tumor in a patient, comprising:
- (a) detecting the level of expression in a tissue sample of two or more genes from Tables 3-9; wherein differential expression of the genes in Tables 3-9 is indicative of metastatic liver cancer.

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- 10. A method of detecting the progression of a metastatic liver cancer in a patient, comprising:
- (a) detecting the level of expression in a tissue sample of two or more genes from Tables 3-9; wherein differential expression of the genes in Tables 3-9 is indicative of a metastatic liver cancer progression.
 - A method of monitoring the treatment of a patient with a metastatic liver cancer, comprising:
 - (a) administering a pharmaceutical composition to the patient;
- 15 (b) preparing a gene expression profile from a cell or tissue sample from the patient; and
 - (c) comparing the patient gene expression profile to a gene expression from a cell population comprising normal liver cells or to a gene expression profile from a cell population comprising metastatic liver tumor cells or to both.

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- 12. A method of treating a patient with a metastatic liver tumor, comprising:
- (a) administering to the patient a pharmaceutical composition, wherein the composition alters the expression of at least one gene in Tables 3-9;
- (b) preparing a gene expression profile from a cell or tissue sample from the patient comprising metastatic liver tumor cells; and
- (c) comparing the patient expression profile to a gene expression profile from an untreated cell population comprising metastatic liver tumor cells.
- 13. A method of differentiating metastatic liver cancer from hepatocellular carcinoma in a patient, comprising:

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- (a) detecting the level of expression in a tissue sample of two or more genes from Tables 3-9; wherein differential expression of the genes in Tables 3-9 is indicative of metastatic liver cancer rather than hepatocellular carcinoma.
- 5 14. A method of screening for an agent capable of modulating the onset or progression of liver cancer, comprising:
 - (a) preparing a first gene expression profile of a cell population comprising liver cancer cells, wherein the expression profile determines the expression level of one or more genes from Tables 3-9;
 - (b) exposing the cell population to the agent;
 - (c) preparing second gene expression profile of the agent-exposed cell population; and
 - (d) comparing the first and second gene expression profiles.
- 15 15. The method of claim 14, wherein the liver cancer is a hepatocellular carcinoma.
 - 16. The method of claim 14, wherein the liver cancer is a metastatic liver cancer.
- 20 17. A composition comprising at least two oligonucleotides, wherein each of the oligonucleotides comprises a sequence that specifically hybridizes to a gene in Tables 3-9.
 - 18. A composition according to claim 17, wherein the composition comprises at least 3 oligonucleotides.
 - 19. A composition according to claim 17, wherein the composition comprises at least 5 oligonucleotides.
- 20. A composition according to claim 17, wherein the composition comprises at 30 least 7 oligonucleotides.

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21. A composition according to claim 17, wherein the composition comprises at least 10 oligonucleotides.

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- 22. A composition according to any one of claims 17-21, wherein the oligonucleotides are attached to a solid support.
 - 23. A composition according to claim 22, wherein the solid support is selected from a group consisting of a membrane, a glass support, a filter, a tissue culture dish, a polymeric material, a bead and a silica support.

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- 24. A solid support comprising at least two oligonucleotides, wherein each of the oligonucleotides comprises a sequence that specifically hybridizes to a gene in Tables 3-9.
- 25. A solid support according to claim 24, wherein the oligonucleotides are covalently attached to the solid support.
 - 26. A solid support according to claim 24, wherein the oligonucleotides are non-covalently attached to the solid support.
- 27. A solid support according to claim 24, wherein the support comprises at least about 10 different oligonucleotides in discrete locations per square centimeter.
 - 28. A solid support according to claim 24, wherein the support comprises at least about 100 different oligonucleotides in discrete locations per square centimeter.

- 29. A solid support according to claim 24, wherein the support comprises at least about 1000 different oligonucleotides in discrete locations per square centimeter.
- 30. A solid support according to claim 24, wherein the support comprises at least about 10,000 different oligonucleotides in discrete locations per square centimeter.
 - 31. A computer system comprising:

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(a) a database containing information identifying the expression level in liver tissue of a set of genes comprising at least two genes in Tables 3-9; and

- (b) a user interface to view the information.
- 5 32. A computer system of claim 31, wherein the database further comprises sequence information for the genes.
 - 33. A computer system of claim 31, wherein the database further comprises information identifying the expression level for the set of genes in normal liver tissue.
 - 34. A computer system of claim 31, wherein the database further comprises information identifying the expression level of the set of genes in liver cancer tissue.
- 35. A computer system of claim 34, wherein the liver cancer tissue comprises hepatocellular carcinoma cells.
 - 36. A computer system of claim 34, wherein the liver cancer tissue comprises metastatic liver cancer cells.
- 20 37. A computer system of any of claims 31-36, further comprising records including descriptive information from an external database, which information correlates said genes to records in the external database.
 - 38. A computer system of claim 37, wherein the external database is GenBank.
 - 39. A method of using a computer system of any one of claims 31-36 to present information identifying the expression level in a tissue or cell of at least one gene in Tables 3-9, comprising:
- (a) comparing the expression level of at least one gene in Tables 3, 4, 5, 6, 8 or 3-9 in the tissue or cell to the level of expression of the gene in the database.

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- 40. A method of claim 39, wherein the expression level of at least two genes are compared.
- 41. A method of claim 39, wherein the expression level of at least five genes are 5 compared.
 - 42. A method of claim 39, wherein the expression level of at least ten genes are compared.
- 10 43. A method of claim 39, further comprising displaying the level of expression of at least one gene in the tissue or cell sample compared to the expression level in liver cancer.

FIGURE 1

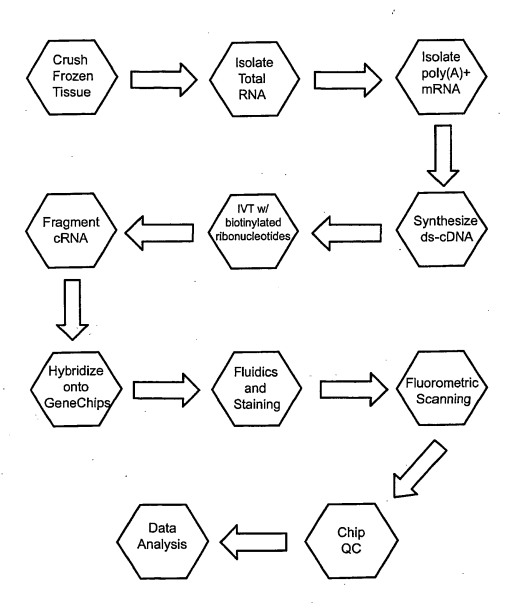


FIGURE 2A

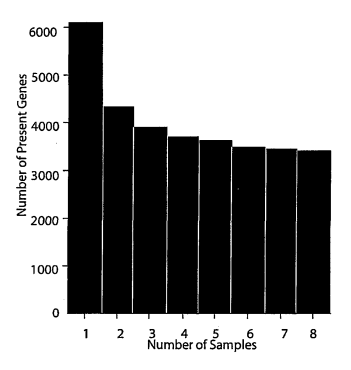


FIGURE 2B

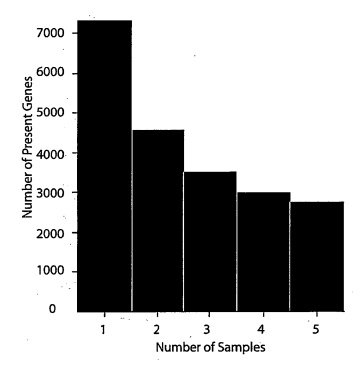
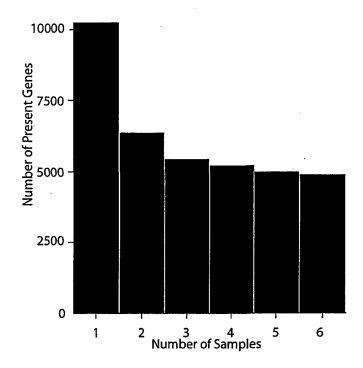


FIGURE 2C



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